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TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C.371				U.S. APPLICATION NUMBER, IF APPLICABLE, 37 CFR 1.5 09752925
INTERNATIONAL APPLICATION PCT/JP98/04475	INTERNATIONAL FILING DATE 05 October 1998 (05.10.98)	PRIORITY DATE CLAIMED 08 October 1997 (08.10.97)		
TITLE OF INVENTION HUMAN PROTEINS HAVING TRANSMEMBRANE DOMAINS AND cDNAs ENCODING THESE PROTEINS				
APPLICANT(S) FOR DO/EO/US Seishi KATO; Tomoko KIMURA; Shingo SEKINE; and Midori KOBAYASHI				
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:				
<ol style="list-style-type: none"> <input checked="" type="checkbox"/> This is a FIRST submission of items concerning a filing under 35 U.S.C.371. <input type="checkbox"/> This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. <input type="checkbox"/> This express request to begin national examination procedures (35 U.S.C. 371 (f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371 (b) and PCT Articles 22 and 39(1). <input checked="" type="checkbox"/> A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date. <input checked="" type="checkbox"/> A copy of the International Application as filed (35 U.S.C. 371(c)(2)) <ol style="list-style-type: none"> <input type="checkbox"/> is transmitted herewith (required only if not transmitted by the International Bureau). <input checked="" type="checkbox"/> has been transmitted by the International Bureau. <input type="checkbox"/> is not required, as the application was filed in the United States Receiving Office (RO/US). <input type="checkbox"/> A translation of the International Application into English (35 U.S.C 371(c)(2)). <input checked="" type="checkbox"/> Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)) <ol style="list-style-type: none"> <input type="checkbox"/> are transmitted herewith (required only if not transmitted by the International Bureau). <input type="checkbox"/> have been transmitted by the International Bureau. <input type="checkbox"/> have not been made; however, the time limit for making such amendments has NOT expired. <input checked="" type="checkbox"/> have not been made and will not be made. <input type="checkbox"/> A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). <input checked="" type="checkbox"/> An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)) (unexecuted (5 sheets)); <input type="checkbox"/> A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). 				
Items 11. to 16. below concern document(s) or information included:				
<ol style="list-style-type: none"> <input type="checkbox"/> An Information Disclosure Statement under 37 CFR 1.97 and 1.98. <input type="checkbox"/> An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included <input type="checkbox"/> A FIRST preliminary amendment. <input type="checkbox"/> A SECOND or SUBSEQUENT preliminary amendment. <input type="checkbox"/> A substitute specification. <input type="checkbox"/> A change of power of attorney and/or address letter. <input checked="" type="checkbox"/> Other items or information: Transmittal Letter (2 sheets in duplicate); PCT Notification of Receipt of Record Copy (Form PCT/IB/301) (1 sheet); PCT Notification Concerning Submission or Transmittal of Priority Document (Form PCT/IB/304) (1 sheet); PCT Notification of the Recording of a Change (Form PCT/IB/306) (1 sheet); PCT Notice Informing the Applicant of the Communication of the International Application to the Designated Offices (Form PCT/IB/308) (1 sheet); PCT Notification of Receipt of Demand by Competent International Preliminary Examining Authority (Form PCT/IEPA/402) (1 sheet); PCT Information Concerning Elected Offices Notified of their Election (Form PCT/IB/332) (1 sheet); PCT International Published Application (WO 99/18203) (without International Search Report) (139 sheets); Cover sheet of PCT International Published Application (WO 99/18203) (with International Search Report attached) (4 sheets); PCT International Preliminary Examination Report (6 sheets); Check (#036342) (\$1100); Certificate of Express Mailing (1 sheet); and Return Postcard. 				

17. The following fees are submitted:**BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)) .(a/o January 1, 2000):**

Search Report has been prepared by the EPO or JPO.....	\$840
International preliminary examination fee paid to USPTO (37 CFR 1.482).....	\$670
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International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(2)-(4).....	\$96

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CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE	
Total claims	11 -20 =	0	X \$18.00	\$--
Independent claims	2 -3 =	0	X \$78.00	\$--
MULTIPLE DEPENDENT CLAIM(S) (if applicable)			+ 260.00	\$260
TOTAL OF ABOVE CALCULATIONS =				\$1100
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TOTAL NATIONAL FEE =				\$1100
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property				\$--
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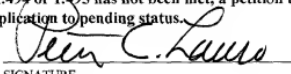
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NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

SEND ALL CORRESPONDENCE TO:

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REGISTRATION NUMBER

528 Rec'd PCT/PTO 07 APR 2000

(b) PRTS

DESCRIPTION

Human Proteins Having Transmembrane
Domains and DNAs Encoding these Proteins

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TECHNICAL FIELD

The present invention relates to human proteins having transmembrane domains and cDNAs coding for these proteins as well as eucaryotic cells expressing said cDNAs. The proteins of the

10 present invention can be employed as pharmaceuticals or as antigens for preparing antibodies against said proteins. The human cDNAs of the present invention can be utilized as probes for the gene diagnosis and gene sources for the gene therapy. Furthermore, the cDNAs can be utilized as gene sources for large-scale production of the proteins encoded by said cDNAs. Cells, wherein these membrane protein genes are introduced and membrane proteins are expressed in large amounts, can be utilized for detection of the corresponding ligands, screening of novel low-molecular pharmaceuticals, and so on.

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BACKGROUND ART

Membrane proteins play important roles, as signal receptors, ion channels, transporters, etc. in the material transportation and the information transmission which are mediated by the cell 25 membrane. Examples thereof include receptors for a variety of cytokines, ion channels for the sodium ion, the potassium ion, the chloride ion, etc., transporters for saccharides and amino

acids, and so on, where the genes of many of them have been cloned already.

It has been clarified that abnormalities of these membrane proteins are associated with a number of hitherto-cryptogenic 5 diseases. For instance, a gene of a membrane protein having twelve transmembrane domains was identified as the gene responsible for cystic fibrosis [Rommens, J. M. et al., Science 245: 1059-1065 (1989)]. In addition, it has been clarified that several membrane proteins act as receptors when a virus infects the cells. For 10 instance, HIV-1 is revealed to infect into the cells through mediation of a membrane protein fusin having a membrane protein on the T-cell membrane, a CD-4 antigen, and seven transmembrane domains [Feng, Y. et al., Science 272: 872-877 (1996)]. Therefore, discovery of a new membrane protein is anticipated to lead to 15 elucidation of the causes of many diseases, so that isolation of a new gene coding for the membrane protein has been desired.

Heretofore, owing to difficulty in the purification, many membrane proteins have been isolated by an approach from the gene side. A general method is the so-called expression cloning which 20 comprises transfection of a cDNA library in eucaryotic cells to express cDNAs and then detection of the cells expressing the target membrane protein on the membrane by an immunological technique using an antibody or a physiological technique on the change in the membrane permeability. However, this method is applicable only 25 to cloning of a gene of a membrane protein with a known function.

In general, membrane proteins possess hydrophobic transmembrane domains inside the proteins, wherein, after

synthesis thereof in the ribosome, these domains remain in the phospholipid membrane to be trapped in the membrane. Accordingly, the evidence of the cDNA for encoding the membrane protein is provided by determination of the whole base sequence of a 5 full-length cDNA followed by detection of highly hydrophobic transmembrane domains in the amino acid sequence of the protein encoded by said cDNA.

DISCLOSURE OF INVENTION

10 The object of the present invention is to provide novel human proteins having transmembrane domains and DNAs coding for said proteins as well as transformation eucaryotic cells that are capable of expressing said cDNAs.

15 As the result of intensive studies, the present inventors have been successful in cloning of cDNAs coding for proteins having transmembrane domains from the human full-length cDNA bank, thereby completing the present invention. In other words, the present invention provides human proteins having transmembrane domains, namely proteins containing any of the amino acid 20 sequences represented by Sequence Nos. 1 to 10. Moreover, the present invention provides DNAs coding for the above-mentioned proteins, exemplified by cDNAs containing any of the base sequences represented by Sequence Nos. 11 to No. 20, as well as transformation eucaryotic cells that are capable of expressing 25 said cDNAs.

BRIEF DESCRIPTION OF DRAWINGS

Figure 1: A figure depicting the -- hydrophobicity/hydrophilicity profile of the protein encoded by clone HP01244.

5 Figure 2: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by clone HP01498.

Figure 3: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by

10 clone HP01565.

Figure 4: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by clone HP01606.

15 Figure 5: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by clone HP01737.

Figure 6: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by clone HP01962.

20 Figure 7: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by clone HP10435.

Figure 8: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by

25 clone HP10479.

Figure 9: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by

clone HP10481.

Figure 10: A figure depicting the hydrophobicity/hydrophilicity profile of the protein encoded by clone HP10495.

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BEST MODE FOR CARRYING OUT OF THE INVENTION

The proteins of the present invention can be obtained, for example, by a method for isolation from human organs, cell lines, etc., a method for preparation of peptides by the chemical

10 synthesis, or a method for production with the recombinant DNA technology using the DNAs coding for the transmembrane domains of the present invention, wherein the method for obtainment by the recombinant DNA technology is employed preferably. For instance, in vitro expression of the proteins can be achieved by preparation of an RNA by in vitro transcription from a vector having one of cDNAs of the present invention, followed by in vitro translation using this RNA as a template. Also, recombination of the translation region into a suitable expression vector by the method known in the art leads to production of a large amount of

15 20 the encoded protein by using prokaryotic cells such as *Escherichia coli*, *Bacillus subtilis*, etc., and eucaryotic cells such as yeasts, insect cells, mammalian cells, etc.

In the case in which a protein of the present invention is produced by a microorganism such as *Escherichia coli* etc., a recombinant expression vector bearing the translation region in the cDNA of the present invention is constructed in an expression vector having an origin, a promoter, a ribosome-binding site, a

cDNA-cloning site, a terminator etc., which can be replicated in the microorganism, and, after transformation of the host cells with said expression vector, the thus-obtained transformant is incubated, whereby the protein encoded by said cDNA can be produced

5 on a large scale in the microorganism. In this case, a protein fragment containing an optional region can be obtained by carrying out the expression with inserting an initiation codon and a termination codon in front of and behind an optional translation region. Alternatively, a fusion protein with another protein can
10 be expressed. Only a protein portion coding for said cDNA can be obtained by cleavage of said fusion protein with a suitable protease.

In the case in which one of the proteins of the present invention is produced in eucaryotic cells, the protein of the present invention can be produced as a transmembrane protein on the cell-membrane surface, when the translation region of said cDNA is subjected to recombination to an expression vector for eucaryotic cells that has a promoter, a splicing region, a poly(A) insertion site, etc., followed by introduction into the eucaryotic cells. The expression vector is exemplified by pKAl, Ped6dpc2, pCDM8, pSVK3, pMSG, pSVL, pBK-CMV, pBK-RSV, EBV vector, pRS, pYES2, and so on. Examples of eucaryotic cells to be used in general include mammalian culture cells such as simian kidney cells COS7, Chinese hamster ovary cells CHO, etc., budding yeasts, fission yeasts, silkworm cells, *Xenopus laevis* egg cells, and so on, but any eucaryotic cells may be used, provided that they are capable of expressing the present proteins on the membrane surface. The

expression vector can be introduced in the eucaryotic cells by methods known in the art such as the electroporation method, the potassium phosphate method, the liposome method, the DEAE-dextran method, and so on.

5 After one of the proteins of the present invention is expressed in prokaryotic cells or eucaryotic cells, the objective protein can be isolated from the culture and purified by a combination of separation procedures known in the art. Such examples include treatment with a denaturing agent such as urea

10 or a surface-active agent, sonication, enzymatic digestion, salting-out or solvent precipitation, dialysis, centrifugation, ultrafiltration, gel filtration, SDS-PAGE, isoelectric focusing, ion-exchange chromatography, hydrophobic chromatography, affinity chromatography, reverse phase chromatography, and so on.

15 The proteins of the present invention include peptide fragments (more than 5 amino acid residues) containing any partial amino acid sequence in the amino acid sequences represented by Sequence Nos. 1. to 10. These peptide fragments can be utilized as antigens for preparation of antibodies. Hereupon, among the 20 proteins of the present invention, those having the signal sequence are secreted in the form of maturation proteins on the surface of the cells, after the signal sequences are removed. Therefore, these maturation proteins shall come within the scope of the present invention. The N-terminal amino acid sequences of 25 the maturation proteins can be easily identified by using the method for the cleavage-site determination in a signal sequence [Japanese Patent Kokai Publication No. 1996-187100]. Furthermore,

some membrane proteins undergo the processing on the cell surface to be converted to the secretory forms. Such proteins or peptides in the secretory forms shall come within the scope of the present invention. When sugar chain-binding sites are present in the amino acid sequences, expression in appropriate eucaryotic cells affords proteins wherein sugar chains are added. Accordingly, such proteins or peptides wherein sugar chains are added shall come within the scope of the present invention.

10 The DNAs of the present invention include all DNAs coding for the above-mentioned proteins. Said DNAs can be obtained by using a method by chemical synthesis, a method by cDNA cloning, and so on.

15 The cDNAs of the present invention can be cloned, for example, from cDNA libraries of the human cell origin. These cDNAs are synthesized by using as templates poly(A)⁺ RNAs extracted from human cells. The human cells may be cells delivered from the human body, for example, by the operation or may be the culture cells.

The cDNAs can be synthesized by using any method selected from the Okayama-Berg method [Okayama, H. and Berg, P., Mol. Cell. Biol.

20 2: 161-170 (1982)], the Gubler-Hoffman method [Gubler, U. and Hoffman, J. Gene 25: 263-269 (1983)], and so on, but it is preferred to use the capping method [Kato, S. et al., Gene 150: 243-250 (1994)], as exemplified in Examples, in order to obtain a full-length clone in an effective manner.

25 The primary selection of one of the cDNAs coding for the human proteins having transmembrane domains is carried out by sequencing of a partial base sequence of a cDNA clone selected

at random from cDNA libraries, sequencing of the amino acid sequence encoded by the base sequence, and recognition of the presence or absence of a hydrophobic site in the resulting N-terminal amino acid sequence region. Next, the secondary selection

5 is carried out by determination of the whole sequence by the sequencing and the protein expression by *in vitro* translation.

Ascertainment of cDNAs of the present invention for encoding the proteins having secretory signal sequences is carried out by using the signal sequence detection method [Yokoyama-Kobayashi, M. et

10 al., Gene 163: 193-196 (1995)]. In other words, the ascertainment for a coding portion of an inserted cDNA fragment to function as a signal sequence is provided by fusing a cDNA fragment coding for the N-terminus of the target protein with a cDNA coding for the protease domain of urokinase and then expressing the resulting

15 cDNA in COS7 cells to detect the urokinase activity in the cell culture medium. On the other hand, in the case in which the urokinase activity is not detectable in the cell medium, the N-terminal region is judged to remain in the membrane.

The cDNAs of the present invention are characterized by containing

20 either of the base sequences represented by Sequence Nos. 11 to 20 or the base sequences represented by Sequence Nos. 21, 23, 25, 27, 29, 31, 33, 35, 37 and 39. Table 1 summarizes the clone number (HP number), the cells affording the cDNA, the total base number of the cDNA, and the number of the amino acid residues of the encoded

25 protein, for each of the cDNAs.

Table 1

	Sequence No.	HP No.	Cell	Number of bases	Number of amino acids
5	1, 11, 21	HP 01244	Stomach Cancer	979	123
	2, 12, 22	HP 01498	Stomach Cancer	1279	220
	3, 13, 23	HP 01565	Stomach Cancer	835	81
	4, 14, 24	HP 01606	Stomach Cancer	1256	301
10	5, 15, 25	HP 01737	Stomach Cancer	1305	383
	6, 16, 26	HP 01962	Liver	899	199
	7, 17, 27	HP 10435	Stomach Cancer	905	229
	8, 18, 28	HP 10479	PMA-U937	841	178
	9, 19, 29	HP 10481	PMA-U937	1451	443
15	10, 20, 30	HP 10495	Stomach Cancer	886	130

Hereupon, the same clones as the cDNAs of the present invention can be easily obtained by screening of the cDNA libraries constructed from the human cell lines and human tissues utilized in the present invention by the use of an oligonucleotide probe synthesized on the basis of the cDNA base sequence described in any of Sequence Nos. 11 to 21, 23, 25, 27, 29, 31, 35, 37 and 39.

In general, the polymorphism due to the individual difference is frequently observed in human genes. Accordingly, any cDNA that is subjected to insertion or deletion of one or plural nucleotides and/or substitution with other nucleotides in Sequence Nos. 11 to 21, 23, 25, 27, 29, 31, 35, 37 and 39 shall come within the scope of the present invention.

In a similar manner, any protein that is formed by these modifications comprising insertion or deletion of one or plural amino acids and/or substitution with other amino acids shall come

within the scope of the present invention, as far as the protein possesses the activity of any protein having the amino acid sequences represented by Sequence Nos. 1 to 10.

The cDNAs of the present invention include cDNA fragments 5 (more than 10 bp) containing any partial base sequence in the base sequences represented by Sequence Nos. 11 to 20 or in the base sequences represented by Sequence Nos. 21, 23, 25, 27, 29, 30, 31, 33, 35 and 39. Also, DNA fragments consisting of a sense chain and an anti-sense chain shall come within this scope. These DNA 10 fragments can be utilized as the probes for the gene diagnosis.

In addition to the activities and uses described above, the polynucleotides and proteins of the present invention may exhibit one or more of the uses or biological activities (including those associated with assays cited herein) identified below. Uses 15 or activities described for proteins of the present invention may be provided by administration or use of such proteins or by administration or use of polynucleotides encoding such proteins (such as, for example, in gene therapies or vectors suitable for introduction of DNA).

20 Research Uses and Utilities

The polynucleotides provided by the present invention can be used by the research community for various purposes. The polynucleotides can be used to express recombinant protein for analysis, characterization or therapeutic use; as markers for 25 tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as

molecular weight markers on Southern gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions; to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to 5 hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; as a probe to "subtract-out" known sequences in the process of discovering other novel polynucleotides; for selecting and making oligomers for attachment to a "gene chip" 10 or other support, including for examination of expression patterns; to raise anti-protein antibodies using DNA immunization techniques; and as an antigen to raise anti-DNA antibodies or elicit another immune response. Where the polynucleotide encodes a protein which binds or potentially binds to another protein (such 15 as, for example, in a receptor-ligand interaction), the polynucleotide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803 (1993)) to identify polynucleotides encoding the other protein with which binding occurs or to identify inhibitors of the binding 20 interaction.

The proteins provided by the present invention can similarly be used in assay to determine biological activity, including in a panel of multiple proteins for high-throughput screening; to raise antibodies or to elicit another immune 25 response; as a reagent (including the labeled reagent) in assays designed to quantitatively determine levels of the protein (or its receptor) in biological fluids; as markers for tissues in which

the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in a disease state); and, of course, to isolate correlative receptors or ligands. Where the protein binds or 5 potentially binds to another protein (such as, for example, in a receptor-ligand interaction), the protein can be used to identify the other protein with which binding occurs or to identify inhibitors of the binding interaction. Proteins involved in these binding interactions can also be used to screen for peptide or 10 small molecule inhibitors or agonists of the binding interaction.

Any or all of these research utilities are capable of being developed into reagent grade or kit format for commercialization as research products.

Methods for performing the uses listed above are well known 15 to those skilled in the art. References disclosing such methods include without limitation "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E.F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic 20 Press, Berger, S.L. and A.R. Kimmel eds., 1987.

Nutritional Uses

Polynucleotides and proteins of the present invention can also be used as nutritional sources or supplements. Such uses include without limitation use as a protein or amino acid 25 supplement, use as a carbon source, use as a nitrogen source and use as a source of carbohydrate. In such cases the protein or polynucleotide of the invention can be added to the feed of a

particular organism or can be administered as a separate solid or liquid preparation, such as in the form of powder, pills, solutions, suspensions or capsules. In the case of microorganisms, the protein or polynucleotide of the invention can be added to 5 the medium in or on which the microorganism is cultured.

Cytokine and Cell Proliferation/Differentiation Activity

A protein of the present invention may exhibit cytokine, cell proliferation (either inducing or inhibiting) or cell differentiation (either inducing or inhibiting) activity or may

10 induce production of other cytokines in certain cell populations. Many protein factors discovered to date, including all known cytokines, have exhibited activity in one or more factor dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cytokine activity. The activity of
15 a protein of the present invention is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+ (preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e and CMK.

20 The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A.M. Kruisbeek, D.H. Margulies,
25 E.M. Shevach, W Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in

Humans); Takai et al., *J. Immunol.* 137:3494-3500, 1986; Bertagnolli et al., *J. Immunol.* 145:1706-1712, 1990; Bertagnolli et al., *Cellular Immunology* 133:327-341, 1991; Bertagnolli, et al., *J. Immunol.* 149:3778-3783, 1992; Bowman et al., *J. Immunol.* 152: 1756-1761, 1994.

Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described in: Polyclonal T cell stimulation, Kruisbeek, A.M. and Shevach, E.M. In *Current Protocols in Immunology*. J.E.e.a. Coligan eds. Vol 1 pp. 3.12.1-3.12.14, John Wiley and Sons, Toronto. 1994; and Measurement of mouse and human Interferony, Schreiber, R.D. In *Current Protocols in Immunology*. J.E.e.a. Coligan eds. Vol 1 pp. 6.8.1-6.8.8, John Wiley and Sons, Toronto. 1994.

10 Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described in: Measurement of Human and Murine Interleukin 2 and Interleukin 4, Bottomly, K., Davis, L.S. and Lipsky, P.E. In *Current Protocols in Immunology*. J.E.e.a. Coligan eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto. 1991; deVries et al., *J. Exp. Med.* 173:1205-1211, 1991; Moreau et al., *Nature* 336:690-692, 1988; Greenberger et al., *Proc. Natl. Acad. Sci. U.S.A.* 80:2931-2938, 1983; Measurement of mouse and human interleukin 6-Nordan, R. In *Current Protocols in Immunology*.

15 20 25 J.E.e.a. Coligan eds. Vol 1 pp. 6.6.1-6.6.5, John Wiley and Sons, Toronto. 1991; Smith et al., *Proc. Natl. Acad. Sci. U.S.A.* 83:1857-1861, 1986; Measurement of human Interleukin 11 - Bennett,

F., Giannotti, J., Clark, S.C. and Turner, K. J. In Current Protocols in Immunology. J.E.e.a. Coligan eds. Vol 1 pp. 6.15.1 — John Wiley and Sons, Toronto. 1991; Measurement of mouse and human Interleukin 9 - Ciarletta, A., Giannotti, J., Clark, S.C. and 5 Turner, K.J. In Current Protocols in Immunology. J.E.e.a. Coligan eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto. 1991.

Assays for T-cell clone responses to antigens (which will identify, among others, proteins that affect APC-T cell interactions as well as direct T-cell effects by measuring 10 proliferation and cytokine production) include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A.M. Kruisbeek, D.H. Margulies, E.M. Shevach, W Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse 15 Lymphocyte Function; Chapter 6, Cytokines and their cellular receptors; Chapter 7, Immunologic studies in Humans); Weinberger et al., Proc. Natl. Acad. Sci. USA 77:6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988.

20 Immune Stimulating or Suppressing Activity

A protein of the present invention may also exhibit immune stimulating or immune suppressing activity, including without limitation the activities for which assays are described herein. A protein may be useful in the treatment of various immune 25 deficiencies and disorders (including severe combined immunodeficiency (SCID)), e.g., in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting

the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by viral (e.g., HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious 5 diseases caused by viral, bacterial, fungal or other infection may be treatable using a protein of the present invention, including infections by HIV, hepatitis viruses, herpesviruses, mycobacteria, Leishmania spp., malaria spp. and various fungal infections such as candidiasis. Of course, in this regard, a 10 protein of the present invention may also be useful where a boost to the immune system generally may be desirable, i.e., in the treatment of cancer.

Autoimmune disorders which may be treated using a protein of the present invention include, for example, connective tissue 15 disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitus, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a 20 protein of the present invention may also be useful in the treatment of allergic reactions and conditions, such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired 25 (including, for example, organ transplantation), may also be treatable using a protein of the present invention.

Using the proteins of the invention it may also be possible to immune responses, in a number of ways. Down regulation may

be in the form of inhibiting or blocking an immune response already in progress or may involve preventing the induction of an immune response. The functions of activated T cells may be inhibited by suppressing T cell responses or by inducing specific tolerance 5 in T cells, or both. Immunosuppression of T cell responses is generally an active, non-antigen-specific, process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or anergy 10 in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after exposure to the tolerizing agent has ceased. Operationally, tolerance can be demonstrated by the lack of a T cell response upon reexposure to specific antigen in the absence of the tolerizing agent.

Down regulating or preventing one or more antigen functions 15 (including without limitation B lymphocyte antigen functions (such as, for example, B7)), e.g., preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell 20 function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a molecule which inhibits or 25 blocks interaction of a B7 lymphocyte antigen with its natural ligand(s) on immune cells (such as a soluble, monomeric form of a peptide having B7-2 activity alone or in conjunction with a

monomeric form of a peptide having an activity of another B lymphocyte antigen (e.g., B7-1, B7-3) or blocking antibody), prior to transplantation can lead to the binding of the molecule to the natural ligand(s) on the immune cells without transmitting the corresponding costimulatory signal. Blocking B lymphocyte antigen function in this manner prevents cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, the lack of costimulation may also be sufficient to anergize the T cells, thereby inducing tolerance in a subject.

10 Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of a combination of B lymphocyte 15 antigens.

The efficacy of particular blocking reagents in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac 20 grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the immunosuppressive effects of CTLA4Ig fusion proteins *in vivo* as described in Lenschow et al., *Science* 257:789-792 (1992) and Turka et al., *Proc. Natl. Acad. Sci USA*, 89:11102-11105 (1992). In addition, murine models 25 of GVHD (see Paul ed., *Fundamental Immunology*, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of blocking B lymphocyte antigen function *in vivo* on the development

of that disease.

Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells

5 that are reactive against self tissue and which promote the production of cytokines and autoantibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms.

Administration of reagents which block costimulation of T cells

10 by disrupting receptor:ligand interactions of B lymphocyte antigens can be used to inhibit T cell activation and prevent production of autoantibodies or T cell-derived cytokines which may be involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive

15 T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases.

Examples include murine experimental autoimmune encephalitis,

20 systemic lupus erythmatosis in MRL/lpr/lpr mice or NZB hybrid mice, murine autoimmune collagen arthritis, diabetes mellitus in NOD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., *Fundamental Immunology*, Raven Press, New York, 1989, pp. 840-856).

25 Upregulation of an antigen function (preferably a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune

responses may be in the form of enhancing an existing immune response or eliciting an initial immune response. For example, enhancing an immune response through stimulating B lymphocyte antigen function may be useful in cases of viral infection. In 5 addition, systemic viral diseases such as influenza, the commoncold, and encephalitis might be alleviated by the administration of stimulatory forms of B lymphocyte antigens systemically.

Alternatively, anti-viral immune responses may be enhanced 10 in an infected patient by removing T cells from the patient, costimulating the T cells in vitro with viral antigen-pulsed APCs either expressing a peptide of the present invention or together with a stimulatory form of a soluble peptide of the present invention and reintroducing the in vitro activated T cells into 15 the patient. Another method of enhancing anti-viral immune responses would be to isolate infected cells from a patient, transfect them with a nucleic acid encoding a protein of the present invention as described herein such that the cells express all or a portion of the protein on their surface, and reintroduce the 20 transfected cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to, and thereby activate, T cells in vivo.

In another application, up regulation or enhancement of 25 antigen function (preferably B lymphocyte antigen function) may be useful in the induction of tumor immunity. Tumor cells (e.g., sarcoma, melanoma, lymphoma, leukemia, neuroblastoma, carcinoma) transfected with a nucleic acid encoding at least one peptide of

the present invention can be administered to a subject to overcome tumor-specific tolerance in the subject. If desired, the tumor cell can be transfected to express a combination of peptides. For example, tumor cells obtained from a patient can be transfected 5 ex vivo with an expression vector directing the expression of a peptide having B7-2-like activity alone, or in conjunction with a peptide having B7-1-like activity and/or B7-3-like activity. The transfected tumor cells are returned to the patient to result in expression of the peptides on the surface of the transfected 10 cell. Alternatively, gene therapy techniques can be used to target a tumor cell for transfection in vivo.

The presence of the peptide of the present invention having the activity of a B lymphocyte antigen(s) on the surface of the tumor cell provides the necessary costimulation signal to T cells 15 to induce a T cell mediated immune response against the transfected tumor cells. In addition, tumor cells which lack MHC class I or MHC class II molecules, or which fail to reexpress sufficient amounts of MHC class I or MHC class II molecules, can be transfected with nucleic acid encoding all or a portion of (e.g., a 20 cytoplasmic-domain truncated portion) of an MHC class I α chain protein and β_2 microglobulin protein or an MHC class II α chain protein and an MHC class II β chain protein to thereby express MHC class I or MHC class II proteins on the cell surface. Expression 25 of the appropriate class I or class II MHC in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an

antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor 5 associated antigens and induce tumor specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumor-specific tolerance in the subject.

The activity of a protein of the invention may, among other 10 means, be measured by the following methods:

Suitable assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A.M. Kruisbeek, D.H. Margulies, E.M. Shevach, W Strober, Pub. Greene Publishing 15 Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., 20 J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., J. Immunol. 137:3494-3500, 1986; Bowman et al., J. Virology 25 61:1992-1998; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, proteins — that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, J. Immunol. 144:3028-3033, 1990; and Assays for B cell function: In vitro antibody production, Mond, J.J. and Brunswick, M. In *Current Protocols in Immunology*. J.E.e.a. Coligan eds. Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto. 1994.

Mixed lymphocyte reaction (MLR) assays (which will identify,

among others, proteins that generate predominantly Th1 and CTL responses) include, without limitation, those described in: *Current Protocols in Immunology*, Ed by J. E. Coligan, A.M. Kruisbeek, D.H. Margulies, E.M. Shevach, W Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify,

among others, proteins expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J. Immunol. 134:536-544, 1995; Inaba et al., Journal of Experimental Medicine 173:549-559, 1991; Macatonia et al., Journal of Immunology 154:5071-5079, 1995; Porgador et al., Journal of Experimental Medicine 182:255-260, 1995; Nair et al., Journal of Virology 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., Journal of Experimental

Medicine 169:1255-1264, 1989; Bhardwaj et al., Journal of Clinical Investigation 94:797-807, 1994; and Inaba et al., Journal of Experimental Medicine 172:631-640, 1990.

Assays for lymphocyte survival/apoptosis (which will

5 identify, among others, proteins that prevent apoptosis after superantigen induction and proteins that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Research 10 10 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, Journal of Immunology 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., International Journal of Oncology 1:639-648, 1992.

Assays for proteins that influence early steps of T-cell

15 commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cellular Immunology 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc. Nat. Acad. Sci. USA 88:7548-7551, 1991.

20 Hematopoiesis Regulating Activity

A protein of the present invention may be useful in regulation of hematopoiesis and, consequently, in the treatment of myeloid or lymphoid cell deficiencies. Even marginal biological activity in support of colony forming cells or of 25 factor-dependent cell lines indicates involvement in regulating hematopoiesis, e.g. in supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other

cytokines, thereby indicating utility, for example, in treating various anemias or for use in conjunction with — irradiation/chemotherapy to stimulate the production of erythroid precursors and/or erythroid cells; in supporting the growth and 5 proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e., traditional CSF activity) useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelo-suppression; in supporting the growth and proliferation of megakaryocytes and consequently of platelets 10 thereby allowing prevention or treatment of various platelet disorders such as thrombocytopenia, and generally for use in place of or complimentary to platelet transfusions; and/or in supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned 15 hematopoietic cells and therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantation, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post 20 irradiation/chemotherapy, either in-vivo or ex-vivo (i.e., in conjunction with bone marrow transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy.

The activity of a protein of the invention may, among other 25 means, be measured by the following methods:

Suitable assays for proliferation and differentiation of various hematopoietic lines are cited above.

Assays for embryonic stem cell differentiation (which will identify, among others, proteins that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson et al. *Cellular Biology* 15:141-151, 1995; Keller et al., *Molecular and Cellular Biology* 13:473-486, 1993; McClanahan et al., *Blood* 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, proteins that regulate lympho-hematopoiesis) include, without limitation, those described in:

5 10 Methylcellulose colony forming assays, Freshney, M.G. In *Culture of Hematopoietic Cells*. R.I. Freshney, et al. eds. Vol pp. 265-268, Wiley-Liss, Inc., New York, NY. 1994; Hirayama et al., *Proc. Natl. Acad. Sci. USA* 89:5907-5911, 1992; Primitive hematopoietic colony forming cells with high proliferative potential, McNiece, I.K. and Briddell, R.A. In *Culture of Hematopoietic Cells*. R.I. Freshney, et al. eds. Vol pp. 23-39, Wiley-Liss, Inc., New York, NY. 1994; Neben et al., *Experimental Hematology* 22:353-359, 1994; Cobblestone area forming cell assay, Ploemacher, R.E. In *Culture of Hematopoietic Cells*. R.I. Freshney, et al. eds. Vol pp. 1-21, Wiley-Liss, Inc., New York, NY. 1994; Long term bone marrow cultures in the presence of stromal cells, Spooncer, E., Dexter, M. and Allen, T. In *Culture of Hematopoietic Cells*. R.I. Freshney, et al. eds. Vol pp. 163-179, Wiley-Liss, Inc., New York, NY. 1994; Long term culture initiating cell assay, Sutherland, H.J. In 15 20 25

20 Culture of Hematopoietic Cells. R.I. Freshney, et al. eds. Vol pp. 139-162, Wiley-Liss, Inc., New York, NY. 1994.

Tissue Growth Activity

A protein of the present invention also may have utility in compositions used for bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as for wound healing and tissue repair and replacement, and in the treatment of burns, 5 incisions and ulcers.

A protein of the present invention, which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Such 10 a preparation employing a protein of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced 15 craniofacial defects, and also is useful in cosmetic plastic surgery.

A protein of this invention may also be used in the treatment of periodontal disease, and in other tooth repair processes. Such 20 agents may provide an environment to attract bone-forming cells, stimulate growth of bone-forming cells or induce differentiation of progenitors of bone-forming cells. A protein of the invention may also be useful in the treatment of osteoporosis or osteoarthritis, such as through stimulation of bone and/or 25 cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes.

Another category of tissue regeneration activity that may

be attributable to the protein of the present invention is tendon/ligament formation. A protein of the present invention, which induces tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally

5 formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue,

10 as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue formation induced by a composition of the present invention contributes to the repair of congenital, trauma induced, or other tendon or

15 ligament defects of other origin, and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments.

The compositions of the present invention may provide an environment to attract tendon or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce

20 differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors *ex vivo* for return *in vivo* to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendinitis, carpal tunnel syndrome and other tendon or ligament

25 defects. The compositions may also include an appropriate matrix and/or sequestering agent as a carrier as is well known in the art.

The protein of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve — and brain tissue, i.e. for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical 5 and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a protein may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nerve injuries, peripheral neuropathy and localized neuropathies, and central nervous system 10 diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular 15 diseases such as stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a protein of the invention.

Proteins of the invention may also be useful to promote better or faster closure of non-healing wounds, including without 20 limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

It is expected that a protein of the present invention may also exhibit activity for generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, 25 intestine, kidney, skin, endothelium), muscle (smooth, skeletal or cardiac) and vascular (including vascular endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part

of the desired effects may be by inhibition or modulation of fibrotic scarring to allow normal tissue to regenerate. A protein of the invention may also exhibit angiogenic activity.

A protein of the present invention may also be useful for 5 gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokine damage.

A protein of the present invention may also be useful for 10 promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for tissue generation activity include, without 15 limitation, those described in: International Patent Publication No. WO95/16035 (bone, cartilage, tendon); International Patent Publication No. WO95/05846 (nerve, neuronal); International Patent Publication No. WO91/07491 (skin, endothelium).

Assays for wound healing activity include, without 20 limitation, those described in: Winter, Epidermal Wound Healing, pps. 71-112 (Maibach, HI and Rovee, DT, eds.), Year Book Medical Publishers, Inc., Chicago, as modified by Eaglstein and Mertz, J. Invest. Dermatol 71:382-84 (1978).

Activin/Inhibin Activity

25 A protein of the present invention may also exhibit activin- or inhibin-related activities. Inhibins are characterized by their ability to inhibit the release of follicle stimulating

hormone (FSH), while activins and are characterized by their ability to stimulate the release of follicle stimulating hormone (FSH). Thus, a protein of the present invention, alone or in heterodimers with a member of the inhibin α family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the protein of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin- β group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, United States Patent 4,798,885. A protein of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as cows, sheep and pigs.

The activity of a protein of the invention may, among other means, be measured by the following methods:

20 Assays for activin/inhibin activity include, without limitation, those described in: Vale et al., Endocrinology 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc Natl Acad Sci USA 83:3091-3095, 1986.

25 Chemoractive/Chemokinetic Activity

A protein of the present invention may have chemotactic or chemokinetic activity (e.g., act as a chemokine) for mammalian

cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. Chemotactic and chemokinetic proteins can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic proteins provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example, attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: *Current Protocols in Immunology*,

Ed by J.E. Coligan, A.M. Kruisbeek, D.H. Margulies, E.M. Shevach,
W. Strober, Pub. Greene Publishing Associates and Wiley-
Interscience (Chapter 6.12, Measurement of alpha and beta
Chemokines 6.12.1-6.12.28; Taub et al. J. Clin. Invest.
5 95:1370-1376, 1995; Lind et al. APMIS 103:140-146, 1995; Muller
et al Eur. J. Immunol. 25: 1744-1748; Gruber et al. J. of Immunol.
152:5860-5867, 1994; Johnston et al. J. of Immunol. 153: 1762-1768,
1994.

Hemostatic and Thrombolytic Activity

10 A protein of the invention may also exhibit hemostatic or
thrombolytic activity. As a result, such a protein is expected
to be useful in treatment of various coagulation disorders
(including hereditary disorders, such as hemophilias) or to
enhance coagulation and other hemostatic events in treating wounds
15 resulting from trauma, surgery or other causes. A protein of the
invention may also be useful for dissolving or inhibiting
formation of thromboses and for treatment and prevention of
conditions resulting therefrom (such as, for example, infarction
of cardiac and central nervous system vessels (e.g., stroke).

20 The activity of a protein of the invention may, among other
means, be measured by the following methods:

Assay for hemostatic and thrombolytic activity include,
without limitation, those described in: Linet et al., J. Clin.
Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res.
25 45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79 (1991);
Schaub, Prostaglandins 35:467-474, 1988.

Receptor/Ligand Activity

A protein of the present invention may also demonstrate activity as receptors, receptor ligands or inhibitors or agonists of receptor/ligand interactions. Examples of such receptors and ligands include, without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell-cell interactions and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses). Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for receptor-ligand activity include without limitation those described in: Current Protocols in Immunology, Ed by J.E. Coligan, A.M. Kruisbeek, D.H. Margulies, E.M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of Cellular Adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J. Immunol. Methods

175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

Anti-Inflammatory Activity

Proteins of the present invention may also exhibit anti-inflammatory activity. The anti-inflammatory activity may 5 be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell-cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by 10 stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Proteins exhibiting such activities can be used to treat inflammatory conditions including chronic or acute conditions), including without limitation inflammation associated with infection (such 15 as septic shock, sepsis or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of 20 cytokines such as TNF or IL-1. Proteins of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material.

Tumor Inhibition Activity

In addition to the activities described above for 25 immunological treatment or prevention of tumors, a protein of the invention may exhibit other anti-tumor activities. A protein may

inhibit tumor growth directly or indirectly (such as, for example, via ADCC). A protein may exhibit its tumor inhibitory activity by acting on tumor tissue or tumor precursor tissue, by inhibiting formation of tissues necessary to support tumor growth (such as, 5 for example, by inhibiting angiogenesis), by causing production of other factors, agents or cell types which inhibit tumor growth, or by suppressing, eliminating or inhibiting factors, agents or cell types which promote tumor growth

Other Activities

10 A protein of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents, including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily 15 characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in bone form or shape); effecting biorhythms or circadian cycles or rhythms; 20 effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, cofactors or other nutritional factors or component(s); effecting behavioral characteristics, including, 25 without limitation, appetite, libido, stress, cognition (including cognitive disorders), depression (including depressive disorders)

and violent behaviors; providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting deficiencies of the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein.

Examples

The present invention is embodied in more detail by the following examples, but this embodiment is not intended to restrict the present invention. The basic operations and the enzyme reactions with regard to the DNA recombination are carried out according to the literature ["Molecular Cloning. A Laboratory Manual", Cold Spring Harbor Laboratory, 1989]. Unless otherwise stated, restrictive enzymes and a variety of modification enzymes to be used were those available from TAKARA SHUZO. The manufacturer's instructions were used for the buffer compositions as well as for the reaction conditions, in each of the enzyme reactions. The cDNA synthesis was carried out according to the literature [Kato, S. et al., Gene 150: 243-250 (1994)].

(1) Preparation of Poly(A)⁺ RNA

The histiocyte lymphoma cell line U937 (ATCC CRL 1593)

stimulated by phorbol ester, tissues of stomach cancer delivered by the operation, and the liver were used for human cells to extract mRNAs. The cell line was incubated by a conventional procedure.

After about 1 g of the human cells was homogenized in 20

5 ml of a 5.5 M guanidinium thiocyanate solution, a total mRNA was prepared according to the literature [Okayama, H. et al., "Method in Enzymology", Vol. 164, Academic Press, 1987]. This was subjected to chromatography on oligo(dT)-cellulose column washed with a 20 mM Tris-hydrochloride buffer solution (pH 7.6), 0.5 M
10 NaCl, and 1 mM EDTA to obtain a poly(A)⁺ RNA according to the above-described literature.

(2) Construction of cDNA Library

Ten micrograms of the above-mentioned poly(A)⁺ RNA were dissolved in a 100 mM Tris-hydrochloride buffer solution (pH 8), 15 one unit of an RNase-free, bacterial alkaline phosphatase was added, and the reaction was run at 37°C for one hour. After the reaction solution was subjected to phenol extraction, followed by ethanol precipitation, the resulting pellet was dissolved in a solution containing 50 mM sodium acetate (pH 6), 1 mM EDTA, 0.1% 20 2-mercaptoethanol, and 0.01% Triton X-100. Thereto was added one unit of a tobacco-origin acid pyrophosphatase (Epicentre Technologies) and a total 100 µl volume of the resulting mixture was reacted at 37°C for one hour. After the reaction solution was subjected to phenol extraction, followed by ethanol precipitation, 25 the resulting pellet was dissolved in water to obtain a solution of a decapped poly(A)⁺ RNA.

The decapped poly(A)⁺ RNA and 3 nmol of a chimeric DNA-

RNA oligonucleotide (5'-dG-dG-dG-dG-dA-dA-dT-dT-dC-dG-dA-G-G-A-3') were dissolved in a solution containing 50 mM Tris-hydrochloride buffer solution (pH 7.5), 0.5 mM ATP, 5 mM MgCl₂, 10 mM 2-mercaptoethanol, and 25% polyethylene glycol, whereto was added 50 units of T4RNA ligase and a total 30 μ l volume of the resulting mixture was reacted at 20°C for 12 hours. After the reaction solution was subjected to phenol extraction, followed by ethanol precipitation, the resulting pellet was dissolved in water to obtain a chimeric-oligo-capped poly(A)' RNA.

10 After digestion of vector pKA1 (Japanese Patent Kokai Publication No. 1992-117292) developed by the present inventors with KpnI, about 60 dT tails were added using a terminal transferase. A vector primer to be used below was prepared by digestion of this product with EcoRV to remove a dT tail at one side.

15 After 6 μ g of the previously-prepared chimeric-oligo-capped poly(A)' RNA was annealed with 1.2 μ g of the vector primer, the resulting product was dissolved in a solution containing 50 mM Tris-hydrochloride buffer solution (pH 8.3), 75 mM KCl, 3 mM MgCl₂, 10 mM dithiothreitol, and 1.25 mM dNTP (dATP + dCTP + dGTP + dTTP), 200 units of a reverse transcriptase (GIBCO-BRL) were added, and the reaction in a total 20 μ l volume was run at 42°C for one hour. After the reaction solution was subjected to phenol extraction, followed by ethanol precipitation, the resulting pellet was dissolved in a solution containing 50 mM Tris-hydrochloride buffer solution (pH 7.5), 100 mM NaCl, 10 mM MgCl₂, and 1 mM dithiothreitol. Thereto were added 100 units of EcoRI and a total 20 μ l volume of the resulting mixture was reacted at

37°C for one hour. After the reaction solution was subjected to phenol extraction, followed by ethanol precipitation, the resulting pellet was dissolved in a solution containing 20 mM Tris-hydrochloride buffer solution (pH 7.5), 100 mM KCl, 4 mM MgCl₂,

5 10 mM (NH₄)₂SO₄, and 50 µg/ml of the bovine serum albumin. Thereto were added 60 units of an *Escherichia coli* DNA ligase and the resulting mixture was reacted at 16°C for 16 hours. To the reaction solution were added 2 µl of 2 mM dNTP, 4 units of *Escherichia coli* DNA polymerase I, and 0.1 unit of *Escherichia coli* RNase H and
10 the resulting mixture was reacted at 12°C for one hour and then at 22°C for one hour.

Next, the cDNA-synthesis reaction solution was used for transformation of *Escherichia coli* DH12S (GIBCO-BRL). The transformation was carried out by the electroporation method. A
15 portion of the transformant was sprayed on the 2xYT agar culture medium containing 100 µg/ml ampicillin and the mixture was incubated at 37°C overnight. A colony formed on the agar medium was picked up at random and inoculated on 2 ml of the 2xYT culture medium containing 100 µg/ml ampicillin. After incubation at 37°C
20 overnight, the culture mixture was centrifuged to separate the mycelia, from which a plasmid DNA was prepared by the alkaline lysis method. The plasmid DNA was subjected to double digestion with EcoRI and NotI, followed by 0.8% agarose gel electrophoresis, to determine the size of the cDNA insert. Furthermore, using the
25 thus-obtained plasmid as a template, the sequence reaction was carried out by using an M13 universal primer labeled with a fluorescent dye and a Taq polymerase (a kit of Applied Biosystems)

and then the product was examined with a fluorescent DNA sequencer (Applied Biosystems) to determine an about 400-bp base sequence at the 5'-terminus of the cDNA. The sequence data were filed as the homo/protein cDNA bank database.

5 (3) Selection of cDNAs Encoding Proteins Having Transmembrane Domains

A base sequence registered in the homo/protein cDNA bank was converted to three frames of amino acid sequences and the presence or absence of an open reading frame (ORF) beginning from

10 the initiation codon was examined. Then, the selection was made for the presence of a signal sequence that is characteristic to a secretory protein at the N-terminus of the portion encoded by the ORF. These clones were sequenced from the both 5' and 3' directions by the use of the deletion method using exonuclease

15 III to determine the whole base sequence. The hydrophobicity/hydrophilicity profiles were obtained for proteins encoded by the ORF by the Kyte-Doolittle method [Kyte, J. & Doolittle, R. F., J. Mol. Biol. 157: 105-132 (1982)] to examine the presence or absence of a hydrophobic region. In the case in 20 which there is a hydrophobic region of a putative transmembrane domain in the amino acid sequence of an encoded protein, this protein was judged as a membrane protein.

(4) Functional Verification of Secretory Signal Sequence or Transmembrane Domains

25 It was verified by the method described in the literature [Yokoyama-Kobayashi, M. et al., Gene 163: 193-196 (1995)] that the N-terminal hydrophobic region in the secretory protein clone

candidate obtained in the above-mentioned steps functions as a secretory signal sequence. First, the plasmid containing the target cDNA was cleaved at an appropriate restriction enzyme site existing at the downstream of the portion expected for encoding 5 the secretory signal sequence. In the case in which this restriction site was a protruding terminus, the site was blunt-ended by the Klenow treatment or treatment with the mung-bean nuclease. Digestion with HindIII was further carried out and a DNA fragment containing the SV40 promoter and a cDNA 10 encoding the secretory signal sequence at the downstream of the promoter was separated by agarose gel electrophoresis. The resulting fragment was inserted between HindIII in pSSD3 (DDBJ/EMBL/GenBank Registration No. AB007632) and a restriction enzyme site selected so as to match with the urokinase-coding frame, 15 thereby constructing a vector expressing a fusion protein of the secretory signal sequence of the target cDNA and the urokinase protease domain.

After *Escherichia coli* (host: JM109) bearing the fusion-protein expression vector was incubated at 37°C for 2 hours 20 in 2 ml of the 2xYT culture medium containing 100 µg/ml of ampicillin, the helper phage M13KO7 (50 µl) was added and the incubation was continued at 37°C overnight. A supernatant separated by centrifugation underwent precipitation with polyethylene glycol to obtain single-stranded phage particles. 25 These particles were suspended in 100 µl of 1 mM Tris-0.1 mM EDTA, pH 8 (TE). Also, there were used as controls suspensions of single-stranded phage particles prepared in the same manner from

pSSD3 and from the vector pKA1-UPA containing a full-length cDNA of urokinase [Yokoyama-Kobayashi, M. et al., Gene 163: 193-196 — (1995)].

The culture cells originating from the simian kidney, COS7,
5 were incubated at 37°C in the presence of 5% CO₂ in the Dulbecco's modified Eagle's culture medium (DMEM) containing 10% fetal calf albumin. Into a 6-well plate (Nunc Inc., 3 cm in the well diameter) were inoculated 1 × 10⁵ COS7 cells and incubation was carried out at 37°C for 22 hours in the presence of 5% CO₂. After the culture
10 medium was removed, the cell surface was washed with a phosphate buffer solution and then washed again with DMEM containing 50 mM Tris-hydrochloric acid (pH 7.5) (TDMEM). To the resulting cells was added a suspension of 1 µl of the single-stranded phage suspension, 0.6 ml of the DMEM culture medium, and 3 µl of
15 TRANSFECTAM™ (IBF Inc.) and the resulting mixture was incubated at 37°C for 3 hours in the presence of 5% CO₂. After the sample solution was removed, the cell surface was washed with TDMEM, 2 ml per well of DMEM containing 10% fetal calf albumin was added, and the incubation was carried out at 37°C for 2 days in the presence
20 of 5% CO₂.

To 10 ml of 50 mM phosphate buffer solution (pH 7.4) containing 2% bovine fibrinogen (Miles Inc.), 0.5% agarose, and 1 mM calcium chloride were added 10 units of human thrombin (Mochida Pharmaceutical Co., Ltd.) and the resulting mixture was solidified
25 in a plate of 9 cm in diameter to prepare a fibrin plate. Ten microliters of the culture supernatant of the transfected COS7 cells were spotted on the fibrin plate, which was incubated at

37°C for 15 hours. In the case in which a clear circle appears on the fibrin plate, it is judged that the cDNA fragment codes for the amino acid sequence functioning as a secretory signal sequence.

On the other hand, in case in which a clear circle is not formed,

5 the cells were washed well, then the fibrin sheet was placed on the cells, and incubation was carried out at 37°C for 15 hours. In case in which a clear portion is formed on the fibrin sheet, it indicates that the urokinase activity was expressed on the cell surface. In other words, the cDNA fragment is judged to code for

10 the transmembrane domains.

(5) Protein Synthesis by In Vitro Translation

The plasmid vector bearing the cDNA of the present invention was used for in vitro transcription/translation with a T_{NT} rabbit reticulocyte lysate kit (Promega). In this case, [³⁵S]methionine was added to label the expression product with a radioisotope. Each of the reactions was carried out according to the protocols attached to the kit. Two micrograms of the plasmid was reacted at 30°C for 90 minutes in a total 25 µl volume of the reaction solution containing 12.5 µl of T_{NT} rabbit reticulocyte lysate, 0.5

20 µl of a buffer solution (attached to kit), 2 µl of an amino acid mixture (methionine-free), 2 µl of [³⁵S]methionine (Amersham) (0.37 MBq/µl), 0.5 µl of T7RNA polymerase, and 20 U of RNasin.

To 3 µl of the resulting reaction solution was added 2 µl of the SDS sampling buffer (125 mM Tris-hydrochloric acid buffer, pH 6.8,

25 120 mM 2-mercaptoethanol, 2% SDS solution, 0.025% bromophenol blue, and 20% glycerol) and the resulting mixture was heated at 95°C for 3 minutes and then subjected to SDS-polyacrylamide gel

electrophoresis. The molecular weight of the translation product was determined by carrying out the autoradiograph.

(6) Expression by COS7

Escherichia coli bearing the expression vector of the 5 protein of the present invention was infected with helper phage M13KO7 and single-stranded phage particles were obtained by the above-mentioned procedure. The thus-obtained phage was used for introducing each expression vector in the culture cells originating from the simian kidney, COS7. After incubation at 37°C

10 for 2 days in the presence of 5% CO₂, the incubation was continued for one hour in the culture medium containing [³⁵S]cystine or [³⁵S]methionine. Collection and dissolution of the cells, followed by subjecting to SDS-PAGE, allowed to observe the presence of a band corresponding to the expression product of each protein, on 15 the membrane fraction which did not exist in the COS7 cells. For instance, the molecular weights of HP01498, HP01565, HP01737, HP010435 and HP010495 were respectively 20 kDa, 13 kDa, 52 kDa, 33 kDa and 20 kDa.

(7) Clone Examples

20 <HP01244> (Sequence Nos. 1, 11, and 21)

Determination of the whole base sequence of the cDNA insert of clone HP01244 obtained from cDNA libraries of human stomach cancer revealed the structure consisting of a 15-bp 5'-nontranslation region, a 372-bp ORF, and a 592-bp 3'-nontranslation region. The ORF codes for a protein consisting of 25 123 amino acid residues and there existed a signal-like sequence at the N-terminus and one transmembrane domain at the C-terminal.

Figure 1 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro translation resulted in formation of a translation product of 14 kDa that was almost consistent with the molecular weight of 12,911 predicted from the ORF.

The search of the protein data base by using the amino acid sequence of the present protein revealed that the protein was analogous to the chicken stem cell antigen 2 (GenBank Accession No. L34554). Table 2 shows the comparison of the amino acid sequence between the human protein of the present invention (HP) and the chicken stem cell antigen 2 (GG). Therein, the marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed a homology of 33.9% in the entire region.

Table 2

HS MKAVL LAL MAGL AI QPGTALLCY SCKAQVSNEDCLOVKNCTQLGEQCWT A--R1 RAVGL

GG-MKAEI[EAVI]AAV[CV]ERAHTI[IC]FSCSDASSNWACLT[P]VKCAE[EE]HCVTTYVGVGIGGK

HS LT-VLSKGCSI NCYDDSDQYYVGKKNTCCDTDLCNASGAHALQPAAAIAL--LPALGL

***** * *** *** *** * *** *

GG SGDSLSKGCGSPVCP SAGINIGIAAASVYCCDSEI CNISGSSSVKASYAVLALGILYVSFVV

25 HS 11 WGP01

1

GG. VI RARE

30 Furthermore, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of

sequences that possessed a homology of 90% or more (for example, Accession No. AA476643) in EST, but many sequences were not distinct and the same ORF as that in the present cDNA was not found. <HP01498> (Sequence Nos. 2, 12, and 23)

5 Determination of the whole base sequence of the cDNA insert of clone HP01498 obtained from cDNA libraries of human stomach cancer revealed the structure consisting of a 227-bp 5'-nontranslation region, a 663-bp ORF, and a 389-bp 3'-nontranslation region. The ORF codes for a protein consisting of
9 10 220 amino acid residues and there existed four transmembrane domains. Figure 2 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro translation resulted in formation of a translation product of 22 kDa that was almost consistent with the
15 molecular weight of 23,318 predicted from the ORF.

The search of the protein data base by using the amino acid sequence of the present protein revealed that the protein was analogous to the rat protein RVP1 (NBRF Accession No. A39484). Table 3 shows the comparison of the amino acid sequence between
20 the human protein of the present invention (HP) and the rat protein RVP1(RN). Therein, the marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed
25 a homology of 81.8% in the entire region. Hereupon, the rat protein had a sequence longer by 60 amino acid residues at the C-terminal side.

Table 3

HS MSMGLEITGTALAVLGWLGTIVCCALPMWRVSFIGSNIITSQNIWEGLWMNCVVQSTGQ
. **. *****. *****. *****. ***. * . *****. *****
5 RN MSMSLEITGTSLAVLGWLCTIVCCALPMWRVSFIGSSIITAQITWEGLWMNC-VQSTGQ
HS MQCKVYDSLLALPQDLQAARALIVVAILLAFAGLLVALVGAQCTNCVQDDTAKAKITIVA
*****. *****. *****. *****. *****. *****. *****
RN MQCKMYDSLLALPQDLQAARALIVVILLAFAGLLVALVGAQCTNCVQDETAKAKITIVA
HS GVLFLLAALLTLVPVWSANTIIIRDYFNPVVPEAQKREMGAGLYVGWAAAALQLLGALL
10 *****. *****. *****. *****. *****. *****. *****
RN GVLFLLAAVLTLVPVWSANTIIIRDYFNPVLPEAQKREMGTCGLYVGWAAAALQLLGALL
HS CCSCPPREKKYTTAKVYVSAPRSTGPGASLGTGYDRKDYY
*****. ***. ***. *****. ***. ***
RN CCSCPPRE-KYAPTKILYSAPRSTGPGTGTAYDRKTTSERPGARTPHHHYQPSMYPT
15

Furthermore, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more (for example, 20 Accession No. H72008) in EST, but many sequences were not distinct and the same ORF as that in the present cDNA was not found.

The rat protein RVPI is one of membrane proteins which are induced by androgen withdrawal and apoptosis in the rat ventral prostate [Briechl, M. M. et al., Mol. Endocrinol. 5: 1381-1388 25 (1991)]. Accordingly, the present protein is considered to play an important role in the signal transduction that is associated with apoptosis.

<HP01565> (Sequence Nos. 3, 13, and 25)

Determination of the whole base sequence of the cDNA insert 30 of clone HP01565 obtained from cDNA libraries of human stomach

cancer revealed the structure consisting of a 62-bp 5'-nontranslation region, a 246-bp ORF, and a 527-bp 3'-nontranslation region. The ORF codes for a protein consisting of 81 amino acid residues and there existed two transmembrane domains.

5 Figure 3 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro translation resulted in formation of a translation product of 10 kDa that was almost consistent with the molecular weight of 9,374 predicted from the ORF.

10 The search of the protein data base using the amino acid sequence of the present protein has revealed the presence of sequences that were analogous to the nematode putative protein F49C12.13 (GenBank Accession No. Z68227). Table 4 shows the comparison of the amino acid sequence between the human protein

15 of the present invention (HP) and the nematode putative protein F49C12.13 (CE). Therein, the marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed

20 a homology of 47.4% in the entire region.

Table 4

HS	MAYHGLTVPLIVMSVFWGFVCGFLVPWFIPKGPNRGVIIWLVTCSVCCYLFWL *. .**. .*.**...** **.*****.* *. .***.**.
5	CE MCNFSYFQLQMGILIPLVSVAFWAIIGFGGPWIVPKGPNRGIIQLMIIIMTAVCCWMFWI HS IAILAQLNPLFGPQLKNETIWLKYHWP ...* *****.***. .**.... .*
	CE MVFLHQLNPLIGPQINVKTIRWISEKWDAPNVI.N

10

Furthermore, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more (for example, Accession No. N57319) in EST, but, since they are partial sequences, it can not be judged whether or not any of these sequences codes for the same protein as the protein of the present invention. <HP01606> (Sequence Nos. 4, 14, and 27)

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Determination of the whole base sequence of the cDNA insert of clone HP01606 obtained from cDNA libraries of human stomach cancer revealed the structure consisting of a 124-bp 5'-nontranslation region, a 906-bp ORF, and a 226-bp 3'-nontranslation region. The ORF codes for a protein consisting of 301 amino acid residues and there existed seven transmembrane domains. Figure 4 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro translation resulted in formation of a translation product of 31 kDa that was almost consistent with the molecular weight of 32,594 predicted from the ORF.

The search of the protein data base using the amino acid

sequence of the present protein has revealed the presence of sequences that were analogous to the nematode putative protein F13H11.9 (GenBank Accession No. AF003389). Table 5 shows the comparison of the amino acid sequence between the human protein 5 of the present invention (HP) and the nematode putative protein F13H11.9 (CE). Therein, the marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed 10 a homology of 45.1% in the region of 195 amino acid residues at the C-terminal side.

Table 5

15	HS	MLALRVARGSGALRGAAWAPGTRPSKRACWALLPPVPCCLGCLAERWRLRPAALGLRL
CE	MIVTSMFR	
HS	PGIGQRNHCSGAGKAAPRPAAGAGAAAEPGGQWGPASTPSLYENPWTIPNMLSMTTRIGL	
	*..... * . *** .. **.	
CE	GIACRCELQLLLTPRRMLRNFSSEQKQSPKIESLPPEERGKYKVA-TIPNAICTARIAA	
20	HS	APVLYGLIIEEDFNIALGVFALAGLTDLLDGFIARNWANQRSLGSLADPLADKILISL
	.**.***. * . * .** *****. *** .. * .***. ***. ***. ***..	
CE	TPLIGLYLVQHNFTPAPFLFTVAGATDLDGFIARNVPGQKSLLGSVLDPVADKLLISTM	
HS	HS YSLTYADLIPVPLTYMIISRDVMLIAAVFYVRYRTLPTPRTLAKYFNPCYATARLKPTF	
 ***. ***. *** .. * .**. *** .. * .***. * .***. *** .. * .**.	
25	CE	FITMTYAGIPLPLTSVILRDICLIGGGFYKRYQVMSPPYSLSRFFNPQVSSMQVVPMT
HS	HS ISKVNTAVQLLILVAASLAAPVFNYADSIY--LQILWCFTAFTTAASAYSYYHYGRKTVQV	
	. **. ***. * . ***. *** .. ***. * * . *** * . *	
CE	CE MSKINTVLQITLVALSLSPPVDFSTGANDVIVGLCITGFTTIYSGLQYASGKAIKKI	

30

Furthermore, the search of the GenBank using the base

sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more (for example, -- Accession No. C16798) in EST, but many sequences were not distinct and the same ORF as that in the present cDNA was not found.

5 <HP01737> (Sequence Nos. 5, 15, and 29)

Determination of the whole base sequence of the cDNA insert of clone HP01737 obtained from cDNA libraries of human stomach cancer revealed the structure consisting of a 21-bp 5'-nontranslation region, a 1152-bp ORF, and a 132-bp 3'-nontranslation region. The ORF codes for a protein consisting of 383 amino acid residues and there existed two transmembrane domains. Figure 5 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro translation resulted in formation of a translation product of 45 kDa that was almost consistent with the molecular weight of 43,222 predicted from the ORF.

The search of the protein data base using the amino acid sequence of the present protein has revealed the presence of sequences that were analogous to the nematode putative protein K09E9.2 (GenBank Accession No. Z79602). Table 6 shows the comparison of the amino acid sequence between the human protein of the present invention (HP) and the nematode putative protein K09E9.2 (CE). Therein, the marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed a homology of 42.2% in the region of 195 amino acid residues at

the C-terminal side.

Table 6

MEALGKLKLFQFDAYPKTLEDFRVKTCGGATVIVSGLLMLLFLSELQYLYTTEVHPELYV
* * .**.****.*.*****. *. **.**. * .*. .*.****. *. *
CE MSLLWSLKHFDAYRKPMDDFRVKTLGGLVTLIATIAVLLIVLETKQFLSTEVLEHLFV
HS D-KSRGDKLKNIIDVLFPHMPACAYLSIDAMDVAQEQLDVEHNLFKQRLDKDGPVSSEA
**. *.*. *. ***. *. * ***. *. *. *. *
CE DSTTSDERVHIEFDITFKLPCNFITVDVMDVSSEAQENINDDIYRLRLDPEGRNISESA
HS ERHGLKVEVTVFDPDSLDPDRCESCYGAEADIKCCNTCEDVREAYRRRGWAFKNPDTI
... *.* .. *. ****. *. *. ****. **. **. *. * ..
CE QKIEINQNKTSVETTDVQEVKCGSCYGAADGI-CNCNTCDWKSAYAVKGWQV-NIEEV
HS EQCRREGFSQKMQEQQNECQCQVYGFLEVVKVAGNFHFAPGKSFQQSHVHVDLQSGFLDN
.*. *. *** ..*. ****. ****. *. *. *. ****.
CE EQCKNDKWWKEFNEHKNECCRVTGTVKAVAGNFHLPAGDPDHQAMRSHVHDLHNLDPVK
HS INMTHYIQHLSFGEDYPGIVNPLDHTNVTAPQASMMFQYFVVKVPTVYMKVDEVLRTNQ
... *. *. ****. ***. ***. *.*. *. ****. ****. * . *. *.
CE FDASHTVNHVSFGKSFPGKNYPLDGKVNTDNRGIMYQYVVKVPTRYDYLDRVDQSHQ
HS FSVTRHEKVANGLLGQGLPGFVLYELSPMMVKLTERHSFTHFLTGVCIAIGGMFTVA
****. *. *.*. *. **. **. *. ..*. *.*. *.
CE FSVTTHHK--DLGRFRQSGLPGFFLQYEFSPLMVQYEEFRQSFASFLVSLCAIVGGVFAMA
HS GLIDSLIYHSARAIQKKIDLGKTT
* . *. ****. *.*. *. ***
CE QLVDITIYHSSRYMKSRIAGGLKT

Furthermore, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more (for example, Accession No. H42261) in EST, but, since they are partial sequences, it can not be judged whether or not any of these sequences codes for the same protein as the protein of the present invention.

<HP01962> (Sequence Nos. 6, 16, and 31)

Determination of the whole base sequence of the cDNA insert of clone HP01962 obtained from cDNA libraries of human liver revealed the structure consisting of a 73-bp 5'-nontranslation 5 region, a 600-bp ORF, and a 226-bp 3'-nontranslation region. The ORF codes for a protein consisting of 199 amino acid residues and there existed at least three transmembrane domains. Figure 6 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro 10 translation resulted in formation of a translation product of 21 kDa that was almost consistent with the molecular weight of 22,134 predicted from the ORF.

The search of the protein data base using the amino acid sequence of the present protein has revealed the presence of 15 sequences that were analogous to a rat phosphatidylethanolamine N-methyltransferase (SWISS-PROT Accession No. Q08388). Table 7 shows the comparison of the amino acid sequence between the human protein of the present invention (HP) and the rat phosphatidylethanolamine N-methyltransferase (RN). Therein, the 20 marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed a homology of 80.8% in the entire region.

Table 7

HS MTRLLGYVDPLDPDFVAVITIFNPNLYVNVARWEHKTQLSRAFGSPYLACYSLSVTI
..***** .*****. **. **. *****. *****. *****. .*

5 RN SWLLGYVDPTEPSFVAVLTIVFNPLFWVVVARWEQRTKQLSRAFGSPYLACYSLGSII
HS LLNLFLRSHCFTQAMLSQPRMESLDTPAAYSLGLALLGLGVVLVLSSFFALGFAGTFLGD
****, *****. ****, **. **. * ***** * . *, ****, ****, *****
RN LLNLFLRSHCFTQAMMSQPKMEGLDSHTIYFLGLALLGWGLVFLSSFYALGFTGTFLGD
HS YFGILKEARVTVFPPNLDNPMYWGSTANYLGWAIMHASPTGLLTVLVALVYVVALFE
10 10 *****. ***. ***. *****. *****. *****. *, ****. *
RN YFGILKEASRVTTFPFSVLNDNPMYWGSTANYLGWALMHASPTGLLTVLVALVYVVALFE
HS EPFTAEIYRQKASGSHKRS
*****. **. ***
RN EPFTAEIYRRKATRLHKRS

Furthermore, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more and contained an initiation codon (for example, Accession No. H83024) in EST, but many sequences were not distinct and the same ORF as that in the present cDNA was not found.

The rat phosphatidylethanolamine N-methyltransferase is a membrane protein which is associated with the biosynthesis of phosphatidylethanolamine [Cui, Z. et al., J. Biol. Chem. 268: 16655-16663 (1993)]. The present protein is considered to be a human homologue of the phosphatidylethanolamine N-methyltransferase and is utilized for the diagnosis and treatment of diseases that are associated with abnormalities of this enzyme.

30 <HP10435> (Sequence Nos. 7, 17, and 23)

Determination of the whole base sequence of the cDNA insert of clone HP10435 obtained from cDNA libraries of human stomach cancer revealed the structure consisting of an 8-bp 5'-nontranslation region, a 690-bp ORF, and a 207-bp 3'-nontranslation region. The ORF codes for a protein consisting of 229 amino acid residues and there existed one transmembrane domain each at the N-terminus and at the C-terminus. Figure 7 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. Introduction of an expression vector, wherein the HindIII-BalI fragment containing a cDNA portion coding for the N-terminal 109 amino acid residues of the present protein was inserted into the HindIII-EcoRV site of pSSD3, into the COS7 cells revealed the urokinase activity on the cell surface to indicate that the present protein remains in the membrane. In vitro translation resulted in formation of a translation product of 24 kDa that was almost consistent with the molecular weight of 24,688 predicted from the ORF.

The search of the protein data base using the amino acid sequence of the present protein has not revealed the presence of any known protein having an analogy. Also, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more and contained an initiation codon (for example, Accession No. H87685) in EST, but many sequences were not distinct and the same ORF as that in the present cDNA was not found.

<HP10479> (Sequence Nos. 8, 18, and 35)

Determination of the whole base sequence of the cDNA insert of clone HP10479 obtained from cDNA libraries of the human lymphoma U937 revealed the structure consisting of a 38-bp 5'-nontranslation region, a 537-bp ORF, and a 266-bp 3'-nontranslation region. The ORF codes for a protein consisting of 178 amino acid residues and there existed a signal-like sequence at the N-terminus and one transmembrane domain at the C-terminus. Figure 8 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein.

10 Introduction of an expression vector, wherein the HindIII-BanII (blunt-ended by treatment with T4DNA polymerase) fragment containing a cDNA portion coding for the N-terminal 45 amino acid residues of the present protein was inserted into the HindIII-SmaI site of pSSD3, into the COS7 cells revealed the urokinase activity 15 in the culture medium to indicate that the present protein is the type-I membrane protein. In vitro translation resulted in formation of a translation product of 33 kDa that was larger than the molecular weight of 19,453 predicted from the ORF. Application of the (-3,-1) rule, a method for predicting the cleavage site 20 in the secretory signal sequence, allows to expect that the maturation protein starts from glutamine at position 22.

The search of the protein data base using the amino acid sequence of the present protein has revealed the presence of sequences that were analogous to the mouse ion channel homologue 25 RIC (GenBank Accession No. U72680). Table 8 shows the comparison of the amino acid sequence between the human protein of the present invention (HP) and the mouse ion channel homologue RIC (MM).

Therein, the marks of -, *, and . represent a gap, an amino acid residue identical with the protein of the present invention, and -- an amino acid residue analogous to the protein of the present invention, respectively. The both proteins possessed a homology 5 of 48.1% in the entire region.

Table 8

10	HS	MSPSGRLCLLTIVGLILPTRGQTLKD TSSSSADSTIMDIQVPTRAPDAVYTELQPTSPT
	MM	MSLSSRLCLLTIVALILPSRGQTPKKPTSIFTADQTSATTRDNVPDPDQTSPGVQTTPLI
	HS	PTWPADETPQ--PQTQTQQLEG-TDGPLVTDPETHKSTKAHPTDDTTLSERPSPSTDV
	**.****. ... *.*..*. * . * ... *. ***..
15	MM	WTREEATGSQTAQQTETQQLTKMATSNPVSDPGPHTSSKKGTP--AVSRIEPLSPSKNF
	HS	QTDPQTLKPSGFHEDDPFFYDEHTLRKRGLLVAAVLFITGIIILTSGKCRQLSRLCRNH
	 * . . * .**. **. ***** ***** ***** ***** . * .
	MM	MPPSYIEHPLDSNENNPFYYDDTLRKRGLLVAAVLFITGIIILTSGKCRQLSQFCLNRH
	HS	R
		*
20	MM	R

Furthermore, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of 25 sequences that possessed a homology of 90% or more (for example, Accession No. AA296696) in EST, but, since they are partial sequences, it can not be judged whether or not any of these sequences codes for the same protein as the protein of the present invention.

30 The mouse ion channel homologue RIC is one of proteins which are induced on introduction of E2a-Pbx1 oncoprotein into the

NIH3T3 fibroblast cells and has been considered to play an important role in the cell cycle and proliferation [Fu, X. et al., Mol. Cell. Biol. 17: 1503-1512 (1997)].

<HP10481> (Sequence Nos. 9, 19, and 37)

5 Determination of the whole base sequence of the cDNA insert of clone HP10481 obtained from cDNA libraries of the human lymphoma U937 revealed the structure consisting of a 104-bp 5'-nontranslation region, a 1332-bp ORF, and a 15-bp 3'-nontranslation region. The ORF codes for a protein consisting of
10 443 amino acid residues and there existed one transmembrane domain at the N-terminus. Figure 9 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. Introduction of an expression vector, wherein the HindIII-PvuII fragment containing
15 a cDNA portion coding for the N-terminal 148 amino acid residues of the present protein was inserted into the HindIII-EcoRV site of pSSD3, into the COS7 cells revealed the urokinase activity on the cell surface to indicate that the present protein is the type-II membrane protein. In vitro translation resulted in formation of
20 a translation product of 51 kDa that was almost consistent with the molecular weight of 51,145 predicted from the ORF.

The search of the protein data base using the amino acid sequence of the present protein has not revealed the presence of any known protein having an analogy. Also, the search of the GenBank 25 using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more (for example, Accession No. AA354554) in EST, but any of the

sequences was shorter than the present cDNAs and was not found to contain the initiation codon.

<HP10495> (Sequence Nos. 10, 20, and 39)

Determination of the whole base sequence of the cDNA insert

5 of clone HP10495 obtained from cDNA libraries of human stomach cancer revealed the structure consisting of a 62-bp 5'-nontranslation region, a 393-bp ORF, and a 431-bp 3'-nontranslation region. The ORF codes for a protein consisting of 130 amino acid residues and there existed two transmembrane
10 domains. Figure 10 depicts the hydrophobicity/hydrophilicity profile, obtained by the Kyte-Doolittle method, of the present protein. In vitro translation resulted in formation of a translation product of 25 kDa that was larger than the molecular weight of 14,964 predicted from the ORF.

15 The search of the protein data base using the amino acid sequence of the present protein has not revealed the presence of any known protein having an analogy. Also, the search of the GenBank using the base sequences of the present cDNA has revealed the presence of sequences that possessed a homology of 90% or more
20 (for example, Accession No. AA431001) in EST, but each of them was shorter than the present cDNA and was not found to contain the initiation codon.

INDUSTRIAL APPLICABILITY

25 The present invention provides human proteins having transmembrane domains and cDNAs coding for these proteins as well as eucaryotic cells expressing said cDNAs. All of the proteins

of the present invention exist in the cell membrane, so that they are considered to be proteins controlling the proliferation and — the differentiation of the cells. Accordingly, the proteins of the present invention can be employed as pharmaceuticals such as 5 carcinostatic agents relating to the control of the proliferation and the differentiation of the cells or as antigens for preparing antibodies against said proteins. The cDNAs of the present invention can be utilized as probes for the gene diagnosis and gene sources for the gene therapy. Furthermore, the cDNAs can be 10 utilized for large-scale expression of said proteins. Cells, wherein these membrane protein genes are introduced and membrane proteins are expressed in large amounts, can be utilized for detection of the corresponding ligands, screening of novel low-molecular pharmaceuticals, and so on.

15 The present invention also provides genes corresponding to the polynucleotide sequences disclosed herein. "Corresponding genes" are the regions of the genome that are transcribed to produce the mRNAs from which cDNA polynucleotide sequences are derived and may include contiguous regions of the 20 genome necessary for the regulated expression of such genes. Corresponding genes may therefore include but are not limited to coding sequences, 5' and 3' untranslated regions, alternatively spliced exons, introns, promoters, enhancers, and silencer or suppressor elements. The corresponding genes can be isolated in 25 accordance with known methods using the sequence information disclosed herein. Such methods include the preparation of probes or primers from the disclosed sequence information for

identification and/or amplification of genes in appropriate genomic libraries or other sources of genomic materials. An "isolated gene" is a gene that has been separated from the adjacent coding sequences, if any, present in the genome of the organism

5 from which the gene was isolated.

Organisms that have enhanced, reduced, or modified expression of the gene(s) corresponding to the polynucleotide sequences disclosed herein are provided. The desired change in gene expression can be achieved through the use of antisense

10 polynucleotides or ribozymes that bind and/or cleave the mRNA transcribed from the gene (Albert and Morris, 1994, Trends Pharmacol. Sci. 15(7): 250-254; Lavarosky et al., 1997, Biochem.

Mol. Med. 62(1): 11-22; and Hampel, 1998, Prog. Nucleic Acid Res. Mol. Biol. 58: 1-39; all of which are incorporated by reference

15 herein). Transgenic animals that have multiple copies of the gene(s) corresponding to the polynucleotide sequences disclosed herein, preferably produced by transformation of cells with genetic constructs that are stably maintained within the transformed cells and their progeny, are provided. Transgenic

20 animals that have modified genetic control regions that increase or reduce gene expression levels, or that change temporal or spatial patterns of gene expression, are also provided (see European Patent No. 0 649 464 B1, incorporated by reference herein).

In addition, organisms are provided in which the gene(s) 25 corresponding to the polynucleotide sequences disclosed herein have been partially or completely inactivated, through insertion of extraneous sequences into the corresponding gene(s) or through

deletion of all or part of the corresponding gene(s). Partial or complete gene inactivation can be accomplished through insertion, preferably followed by imprecise excision, of transposable elements (Plasterk, 1992, *Bioessays* 14(9): 629-633;

5 Zwaal et al., 1993, *Proc. Natl. Acad. Sci. USA* 90(16): 7431-7435; Clark et al., 1994, *Proc. Natl. Acad. Sci. USA* 91(2): 719-722; all of which are incorporated by reference herein), or through homologous recombination, preferably detected by positive/negative genetic selection strategies (Mansour et al.,
10 1988, *Nature* 336: 348-352; U.S. Patent Nos. 5,464,764; 5,487,992; 5,627,059; 5,631,153; 5,614, 396; 5,616,491; and 5,679,523; all of which are incorporated by reference herein). These organisms with altered gene expression are preferably eukaryotes and more preferably are mammals. Such organisms are useful for the
15 development of non-human models for the study of disorders involving the corresponding gene(s), and for the development of assay systems for the identification of molecules that interact with the protein product(s) of the corresponding gene(s).

Where the protein of the present invention is membrane-bound (e.g., is a receptor), the present invention also provides for soluble forms of such protein. In such forms part or all of the intracellular and transmembrane domains of the protein are deleted such that the protein is fully secreted from the cell in which it is expressed. The intracellular and
25 transmembrane domains of proteins of the invention can be identified in accordance with known techniques for determination of such domains from sequence information.

Proteins and protein fragments of the present invention include proteins with amino acid sequence lengths that are at least 25% (more preferably at least 50%, and most preferably at least 75%) of the length of a disclosed protein and have at least 60% sequence identity (more preferably, at least 75% identity; most preferably at least 90% or 95% identity) with that disclosed protein, where sequence identity is determined by comparing the amino acid sequences of the proteins when aligned so as to maximize overlap and identity while minimizing sequence gaps. Also included in the present invention are proteins and protein fragments that contain a segment preferably comprising 8 or more (more preferably 20 or more, most preferably 30 or more) contiguous amino acids that shares at least 75% sequence identity (more preferably, at least 85% identity; most preferably at least 95% identity) with any such segment of any of the disclosed proteins.

Species homologs of the disclosed polynucleotides and proteins are also provided by the present invention. As used herein, a "species homologue" is a protein or polynucleotide with a different species of origin from that of a given protein or polynucleotide, but with significant sequence similarity to the given protein or polynucleotide, as determined by those of skill in the art. Species homologs may be isolated and identified by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source from the desired species.

The invention also encompasses allelic variants of the disclosed polynucleotides or proteins; that is, naturally-

occurring alternative forms of the isolated polynucleotide which also encode proteins which are identical, homologous, or related to that encoded by the polynucleotides.

The invention also includes polynucleotides with sequences 5 complementary to those of the polynucleotides disclosed herein.

The present invention also includes polynucleotides capable of hybridizing under reduced stringency conditions, more preferably stringent conditions, and most preferably highly stringent conditions, to polynucleotides described herein.

10 Examples of stringency conditions are shown in the table below: highly stringent conditions are those that are at least as stringent as, for example, conditions A-F; stringent conditions are at least as stringent as, for example, conditions G-L; and reduced stringency conditions are at least as stringent as, for 15 example, conditions M-R.

Table 9

Stringency Condition	Polynucleotide Hybrid	Hybrid Length (bp) ^f	Hybridization Temperature and Buffer ^f	Wash Temperature and Buffer ^f
A	DNA : DNA	≥50	65°C; 1×SSC -or- 42°C; 1×SSC.50% formamide	65°C; 0.3×SSC
B	DNA : DNA	<50	T _B *: 1×SSC	T _B *: 1×SSC
C	DNA : RNA	≥50	67°C: 1×SSC -or- 45°C: 1×SSC.50% formamide	67°C: 0.3×SSC
D	DNA : RNA	<50	T _D *: 1×SSC	T _D *: 1×SSC
E	RNA : RNA	≥50	70°C: 1×SSC -or- 50°C: 1×SSC.50% formamide	70°C: 0.3×SSC
F	RNA : RNA	<50	T _F *: 1×SSC	T _F *: 1×SSC
G	DNA : DNA	≥50	65°C: 4×SSC -or- 42°C: 4×SSC.50% formamide	65°C: 1×SSC
H	DNA : DNA	<50	T _H *: 4×SSC	T _H *: 4×SSC
I	DNA : RNA	≥50	67°C: 4×SSC -or- 45°C: 4×SSC.50% formamide	67°C: 1×SSC
J	DNA : RNA	<50	T _J *: 4×SSC	T _J *: 4×SSC
K	RNA : RNA	≥50	70°C: 4×SSC -or- 50°C: 4×SSC.50% formamide	67°C: 1×SSC
L	RNA : RNA	<50	T _L *: 2×SSC	T _L *: 2×SSC
M	DNA : DNA	≥50	50°C: 4×SSC -or- 40°C: 6×SSC.50% formamide	50°C: 2×SSC
N	DNA : DNA	<50	T _N *: 6×SSC	T _N *: 6×SSC
O	DNA : RNA	≥50	55°C: 4×SSC -or- 42°C: 6×SSC.50% formamide	55°C: 2×SSC
P	DNA : RNA	<50	T _P *: 6×SSC	T _P *: 6×SSC
Q	RNA : RNA	≥50	60°C: 4×SSC -or- 45°C: 6×SSC.50% formamide	60°C: 2×SSC
R	RNA : RNA	<50	T _R *: 4×SSC	T _R *: 4×SSC

‡ : The hybrid length is that anticipated for the hybridized region(s) of the hybridizing polynucleotides. When hybridizing a polynucleotide to a target polynucleotide of unknown sequence, the hybrid length is assumed to be that of the hybridizing polynucleotide. When polynucleotides of known sequence are hybridized, the hybrid length can be determined by aligning the sequences of the polynucleotides and identifying the region or regions of optimal sequence complementarity.

5 10 † : SSPE (1×SSPE is 0.15M NaCl, 10mM Na₂PO₄, and 1.25mM EDTA, pH7.4) can be substituted for SSC (1×SSC is 0.15M NaCl and 15mM sodium citrate) in the hybridization and wash buffers: washes are performed for 15 minutes after hybridization is complete.

*T_B - T_F : The hybridization temperature for hybrids anticipated to be less than 50 base pairs in length should be 5-10°C less than the melting temperature (T_m) of the hybrid, where T_m is determined according to the following equations. For hybrids less than 18 base pairs in length, T_m(°C)=2(# of A + T bases) + 4(# of G + C bases). For hybrids between 18 and 49 base pairs in length, T_m(°C)=81.5 + 16.6(log₁₀[Na⁺]) + 0.41 (%G+C) - (600/N), where N is the number of bases in the hybrid, and [Na⁺] is the concentration of sodium ions in the hybridization buffer ([Na⁺] for 1×SSC=0.165M).

Additional examples of stringency conditions for polynucleotide hybridization are provided in Sambrook, J., E.F. Fritsch, and T. Maniatis, 1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, 5 NY, chapters 9 and 11, and Current Protocols in Molecular Biology, 1995, F.M. Ausubel et al., eds., John Wiley & Sons, Inc., sections 2.10 and 6.3-6.4, incorporated herein by reference.

Preferably, each such hybridizing polynucleotide has a length that is at least 25% (more preferably at least 50%, and most

10 preferably at least 75%) of the length of the polynucleotide of the present invention to which it hybridizes, and has at least 60% sequence identity (more preferably, at least 75% identity; most preferably at least 90% or 95% identity) with the polynucleotide of the present invention to which it hybridizes, 15 where sequence identity is determined by comparing the sequences of the hybridizing polynucleotides when aligned so as to maximize overlap and identity while minimizing sequence gaps.

CLAIMS

1. A protein comprising any of the amino acid sequences represented by Sequence Nos. 1 to 10.

5 2. A DNA coding for the protein according to Claim 1.

3. A cDNA comprising any of the base sequences represented by Sequence Nos. 11 to 20.

4. The cDNA according to Claim 3 comprising any of the base sequences represented by Sequence Nos. 21, 23, 25, 27, 29,

5 10 31, 33, 35, 37 and 39.

5. A vector capable of expressing the DNA or cDNA according to any of Claims 2 to 4 in in vitro translation or an eucaryotic cell.

6. A transformation eucaryotic cell capable of expressing the DNA or cDNA according to any of Claim 2 to 4 to produce the protein according to Claim 1.



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(54) Title: HUMAN PROTEINS HAVING TRANSMEMBRANE DOMAINS AND cDNAs ENCODING THESE PROTEINS

(55) Abstract

The invention provides human proteins having transmembrane domains and cDNAs coding for these proteins as well as eukaryotic cells expressing said cDNAs. All of the proteins exist in the cell membrane, so that they are considered to be proteins controlling the proliferation and the differentiation of the cells. Accordingly, the proteins can be employed as pharmaceuticals such as carcinostatic agents relating to the control of the proliferation and the differentiation of the cells or as antigens for preparing antibodies against said proteins. The cDNAs can be utilized as probes for the gene diagnosis and gene sources for the gene therapy. Furthermore, the cDNAs can be utilized for large-scale expression of said proteins. Cells, wherein these membrane protein genes are introduced and membrane proteins are expressed in large amounts, can be utilized for detection of the corresponding ligands, screening of novel low-molecular pharmaceuticals, and so on.

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CN	China	KZ	Kazakhstan	RO	Romania		
CU	Cuba	LC	Saint Lucia	RU	Russian Federation		
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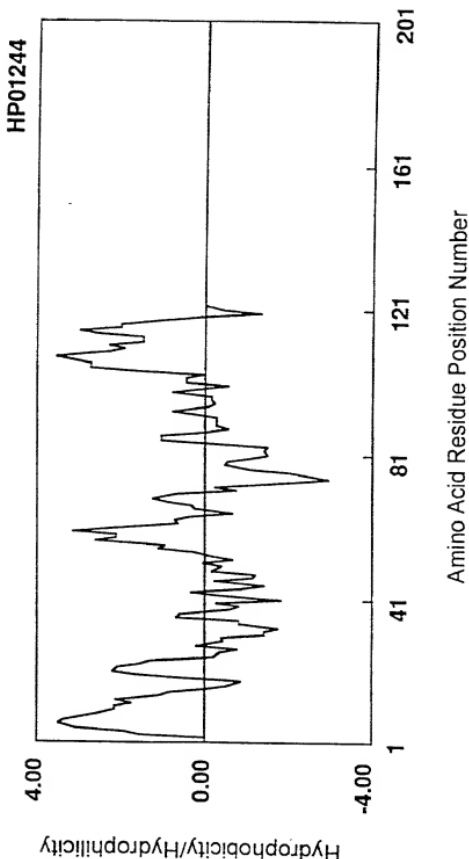


Fig. 1

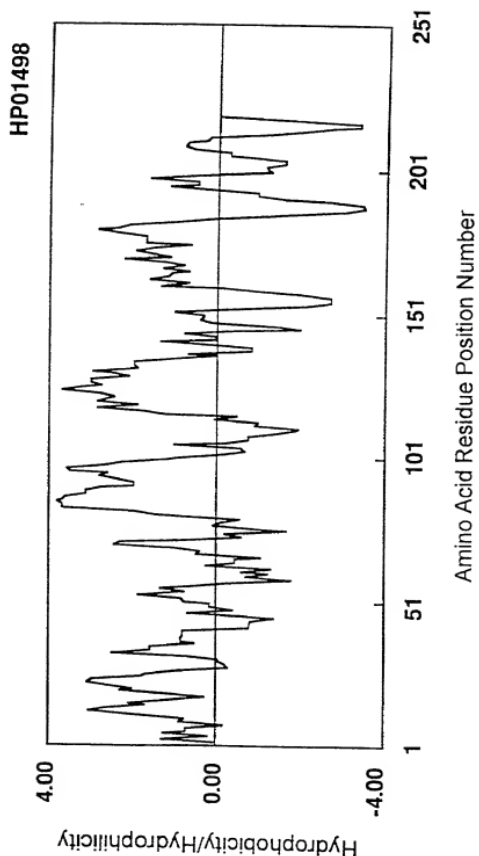


Fig. 2

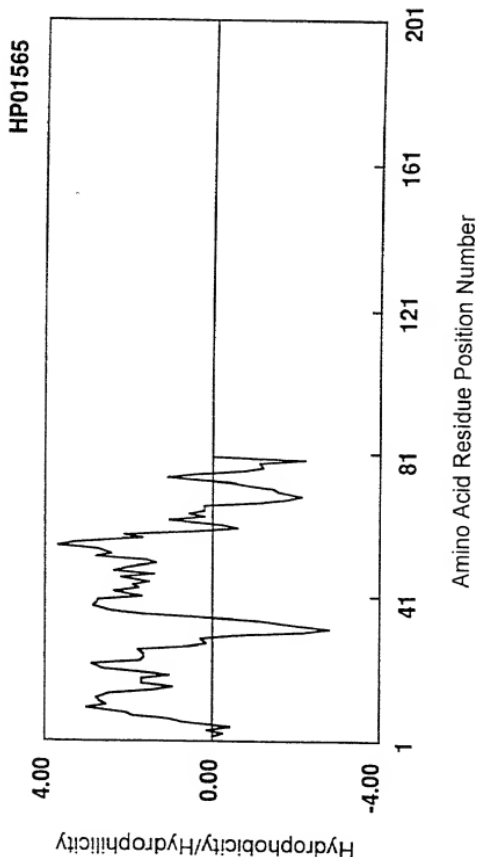


Fig. 3

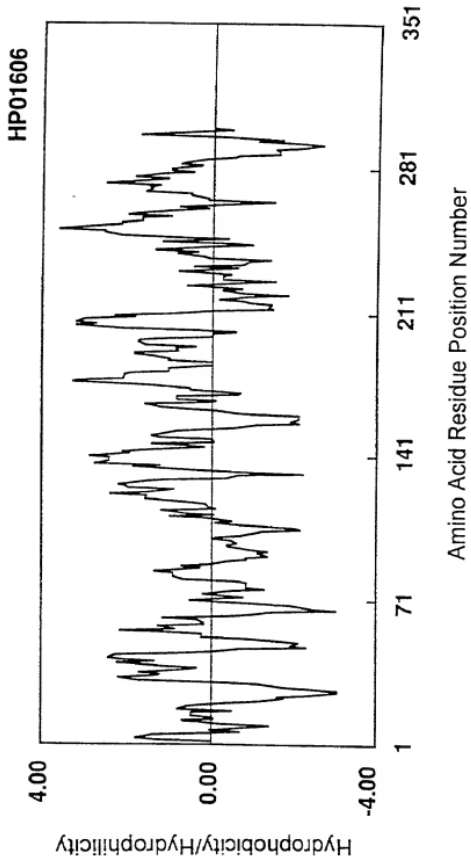


Fig. 4

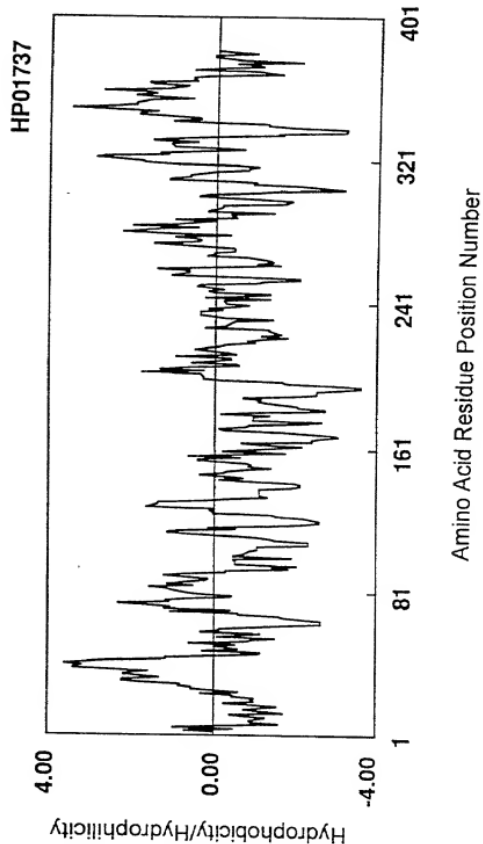


Fig. 5

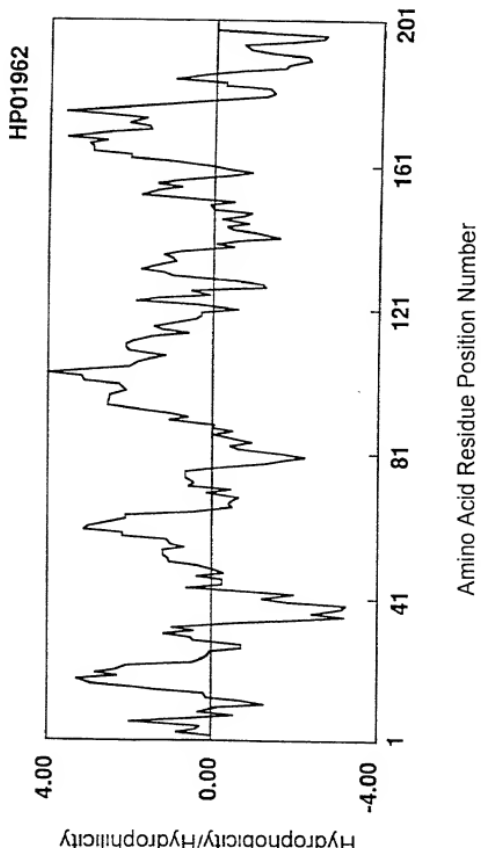


Fig. 6

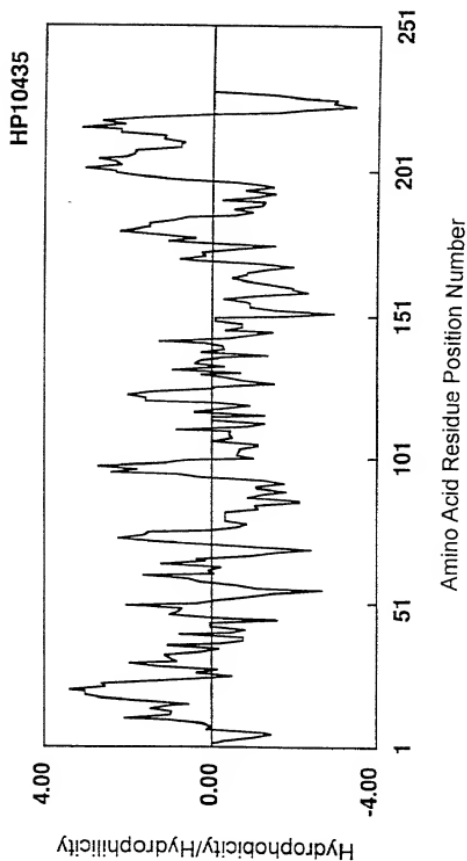


Fig. 7

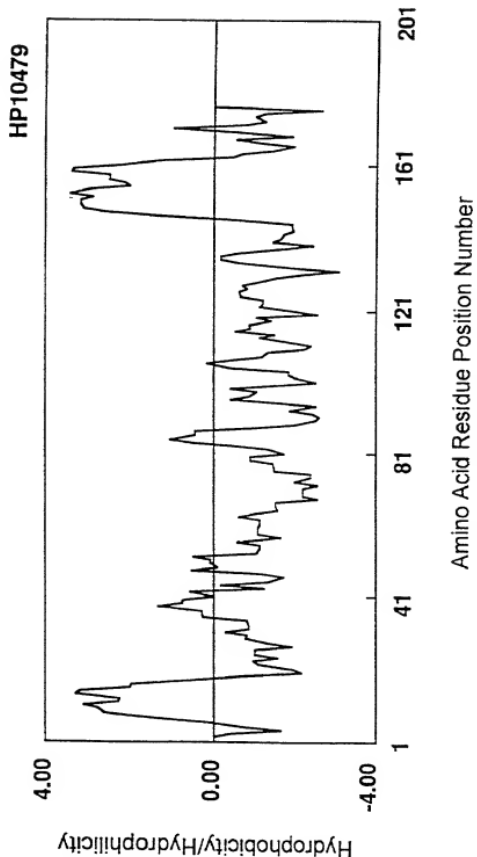


Fig. 8

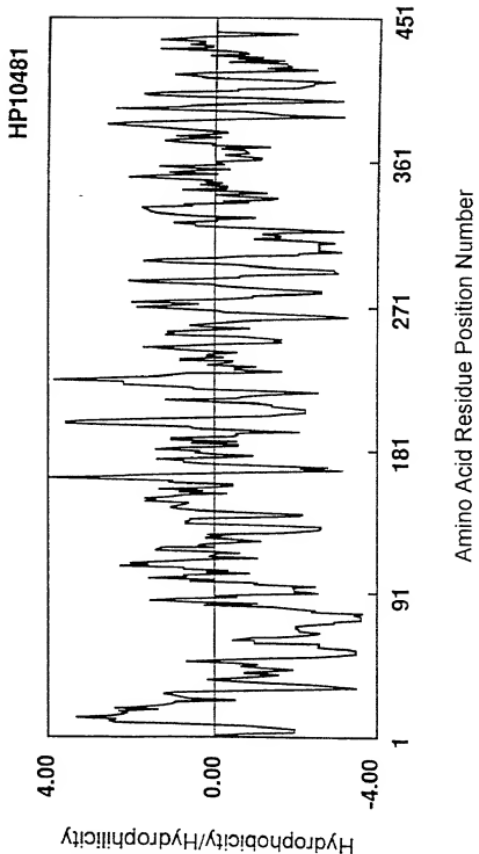


Fig. 9

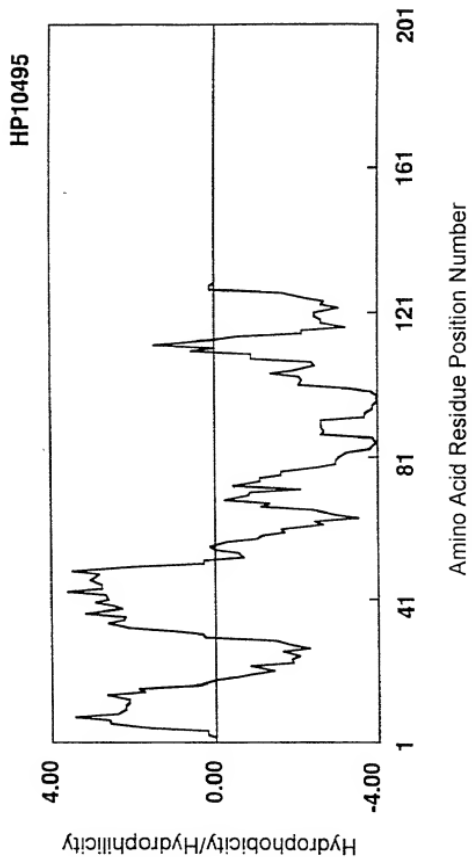


Fig. 10

09/529205

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Trp Thr Ala Arg Ile Arg Ala Val Gly Leu Leu Thr Val Ile Ser Lys

50 55 60

5 Gly Cys Ser Leu Asn Cys Val Asp Asp Ser Gln Asp Tyr Tyr Val Gly

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35 40 45

Trp Arg Leu Arg Pro Ala Ala Leu Gly Leu Arg Leu Pro Gly Ile Gly

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Gln Arg Asn His Cys Ser Gly Ala Gly Lys Ala Ala Pro Arg Pro Ala

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145 150 155 160

15 Arg Ser Ala Leu Gly Ser Ala Leu Asp Pro Leu Ala Asp Lys Ile Leu

165 170 175

Ile Ser Ile Leu Tyr Val Ser Leu Thr Tyr Ala Asp Leu Ile Pro Val

180 185 190

Pro Leu Thr Tyr Met Ile Ile Ser Arg Asp Val Met Leu Ile Ala Ala

20 195 200 205

Val Phe Tyr Val Arg Tyr Arg Thr Leu Pro Thr Pro Arg Thr Leu Ala

210 215 220

Lys Tyr Phe Asn Pro Cys Tyr Ala Thr Ala Arg Leu Lys Pro Thr Phe

225 230 235 240

25 Ile Ser Lys Val Asn Thr Ala Val Gln Leu Ile Leu Val Ala Ala Ser

245 250 255

Leu Ala Ala Pro Val Phe Asn Tyr Ala Asp Ser Ile Tyr Leu Gln Ile

260

265

270

Leu Trp Cys Phe Thr Ala Phe Thr Thr Ala Ala Ser Ala Tyr Ser Tyr

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Val Ser Gly Leu Leu Met Leu Leu Leu Phe Leu Ser Glu Leu Gln Tyr

35 40 45

20 Tyr Leu Thr Thr Glu Val His Pro Glu Leu Tyr Val Asp Lys Ser Arg

50 55 60

Gly Asp Lys Leu Lys Ile Asn Ile Asp Val Leu Phe Pro His Met Pro

65 70 75 80

Cys Ala Tyr Leu Ser Ile Asp Ala Met Asp Val Ala Gly Glu Gln Gln

25 85 90 95

Leu Asp Val Glu His Asn Leu Phe Lys Gln Arg Leu Asp Lys Asp Gly

100 105 110

Ile Pro Val Ser Ser Glu Ala Glu Arg His Glu Leu Gly Lys Val Glu

115 120 125

Val Thr Val Phe Asp Pro Asp Ser Leu Asp Pro Asp Arg Cys Glu Ser

130 135 140

5 Cys Tyr Gly Ala Glu Ala Glu Asp Ile Lys Cys Cys Asn Thr Cys Glu

145 150 155 160

Asp Val Arg Glu Ala Tyr Arg Arg Gly Trp Ala Phe Lys Asn Pro

165 170 175

Asp Thr Ile Glu Gln Cys Arg Arg Glu Gly Phe Ser Gln Lys Met Gln

10 180 185 190

Glu Gln Lys Asn Glu Gly Cys Gln Val Tyr Gly Phe Leu Glu Val Asn

195 200 205

Lys Val Ala Gly Asn Phe His Phe Ala Pro Gly Lys Ser Phe Gln Gln

210 215 220

15 Ser His Val His Val His Asp Leu Gln Ser Phe Gly Leu Asp Asn Ile

225 230 235 240

Asn Met Thr His Tyr Ile Gln His Leu Ser Phe Gly Glu Asp Tyr Pro

245 250 255

Gly Ile Val Asn Pro Leu Asp His Thr Asn Val Thr Ala Pro Gln Ala

20 260 265 270

Ser Met Met Phe Gln Tyr Phe Val Lys Val Val Pro Thr Val Tyr Met

275 280 285

Lys Val Asp Gly Glu Val Leu Arg Thr Asn Gln Phe Ser Val Thr Arg

290 295 300

25 His Glu Lys Val Ala Asn Gly Leu Leu Gly Asp Gln Gly Leu Pro Gly

305 310 315 320

Val Phe Val Leu Tyr Glu Leu Ser Pro Met Met Val Lys Leu Thr Glu

325

330

335

Lys His Arg Ser Phe Thr His Phe Leu Thr Gly Val Cys Ala Ile Ile

340

345

350

Gly Gly Met Phe Thr Val Ala Gly Leu Ile Asp Ser Leu Ile Tyr His

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Ser Ala Arg Ala Ile Gln Lys Lys Ile Asp Leu Gly Lys Thr Thr

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35 40 45

Pro Tyr Leu Ala Cys Tyr Ser Leu Ser Val Thr Ile Leu Leu Leu Asn

50 55 60

Phe Leu Arg Ser His Cys Phe Thr Gln Ala Met Leu Ser Gln Pro Arg

25 65 70 75 80

Met Glu Ser Leu Asp Thr Pro Ala Ala Tyr Ser Leu Gly Leu Ala Leu

85 90 95

Leu Gly Leu Gly Val Val Leu Val Leu Ser Ser Phe Phe Ala Leu Gly

100 105 110

Phe Ala Gly Thr Phe Leu Gly Asp Tyr Phe Gly Ile Leu Lys Glu Ala

115 120 125

5 Arg Val Thr Val Phe Pro Phe Asn Ile Leu Asp Asn Pro Met Tyr Trp

130 135 140

Gly Ser Thr Ala Asn Tyr Leu Gly Trp Ala Ile Met His Ala Ser Pro

145 150 155 160

Thr Gly Leu Leu Leu Thr Val Leu Val Ala Leu Thr Tyr Ile Val Ala

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Ser Gly Ser His Lys Arg Ser

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20 <213> Homo sapiens

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20 25 30

Glu Ile Cys Thr Gln Cys Pro Gly Ser Val Gln Asn Leu Ser Lys Val

35 40 45
Ala Phe Tyr Cys Lys Thr Thr Arg Glu Leu Met Leu His Ala Arg Cys ~
50 55 60
Cys Leu Asn Gln Lys Gly Thr Ile Leu Gly Leu Asp Leu Gln Asn Cys
5 65 70 75 80
Ser Leu Glu Asp Pro Gly Pro Asn Phe His Gln Ala His Thr Thr Val
85 90 95
Ile Ile Asp Leu Gln Ala Asn Pro Leu Lys Gly Asp Leu Ala Asn Thr
100 105 110
10 Phe Arg Gly Phe Thr Gln Leu Gln Thr Leu Ile Leu Pro Gln His Val
115 120 125
Asn Cys Pro Gly Gly Ile Asn Ala Trp Asn Thr Ile Thr Ser Tyr Ile
130 135 140
Asp Asn Gln Ile Cys Gln Gly Gln Lys Asn Leu Cys Asn Asn Thr Gly
15 145 150 155 160
Asp Pro Glu Met Cys Pro Glu Asn Gly Ser Cys Val Pro Asp Gly Pro
165 170 175
Gly Leu Leu Gln Cys Val Cys Ala Asp Gly Phe His Gly Tyr Lys Cys
180 185 190
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Lys Ala Lys Thr Ser
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35 40 45

Ala Val Tyr Thr Glu Leu Gln Pro Thr Ser Pro Thr Pro Trp Pro

50 55 60

15 Ala Asp Glu Thr Pro Gln Pro Gln Thr Gln Gln Leu Glu Gly

65 70 75 80

Thr Asp Gly Pro Leu Val Thr Asp Pro Glu Thr His Lys Ser Thr Lys

85 90 95

Ala Ala His Pro Thr Asp Asp Thr Thr Leu Ser Glu Arg Pro Ser

20 100 105 110

Pro Ser Thr Asp Val Gln Thr Asp Pro Gln Thr Leu Lys Pro Ser Gly

115 120 125

Phe His Glu Asp Asp Pro Phe Phe Tyr Asp Glu His Thr Leu Arg Lys

130 135 140

25 Arg Gly Leu Leu Val Ala Ala Val Leu Phe Ile Thr Gly Ile Ile Ile

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175

Cys Arg

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25

30

15 Arg Gln Ala Pro Ala Gly Ser Pro Arg Gly Leu Arg Lys Gly Ala Ala

35

40

45

Pro Ala Arg Glu Arg Arg Gly Arg Glu Gln Ser Thr Leu Glu Ser Glu

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55

60

Glu Trp Asn Pro Trp Glu Gly Asp Glu Lys Asn Glu Gln Gln His Arg

20 65 70 75 80

Phe Lys Thr Ser Leu Gln Ile Leu Asp Lys Ser Thr Lys Gly Lys Thr

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95

Asp Leu Ser Val Gln Ile Trp Gly Lys Ala Ala Ile Gly Leu Tyr Leu

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105

110

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Gln Trp Arg Glu Gly Lys Ser Ile Val Gly Arg Thr Gln Tyr Ser Phe

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Asn Val Val Leu Ile Leu Asn Gly Arg Glu Lys Ala Lys Ile Phe Tyr

5 165 170 175

Ala Thr Gln Trp Leu Leu Tyr Ala Gln Asn Leu Val Gln Ile Gln Lys

180 185 190

Leu Gln His Leu Ala Val Val Leu Leu Gly Asn Glu His Cys Asp Asn

195 200 205

10 Glu Trp Ile Asn Pro Phe Leu Lys Arg Asn Gly Gly Phe Val Glu Leu

210 215 220

Leu Phe Ile Ile Tyr Asp Ser Pro Trp Ile Asn Asp Val Asp Val Phe

225 230 235 240

Gln Trp Pro Leu Gly Val Ala Thr Tyr Arg Asn Phe Pro Val Val Glu

15 245 250 255

Ala Ser Trp Ser Met Leu His Asp Glu Arg Pro Tyr Leu Cys Asn Phe

260 265 270

Leu Gly Thr Ile Tyr Glu Asn Ser Ser Arg Gin Ala Leu Met Asn Ile

275 280 285

20 Leu Lys Lys Asp Gly Asn Asp Lys Leu Cys Trp Val Ser Ala Arg Glu

290 295 300

His Trp Gln Pro Gln Glu Thr Asn Glu Ser Leu Lys Asn Tyr Gln Asp

305 310 315 320

Ala Leu Leu Gln Ser Asp Leu Thr Leu Cys Pro Val Gly Val Asn Thr

25 325 330 335

Glu Cys Tyr Arg Ile Tyr Glu Ala Cys Ser Tyr Gly Ser Ile Pro Val

340 345 350

Val Glu Asp Val Met Thr Ala Gly Asn Cys Gly Asn Thr Ser Val His

355 360 365

His Gly Ala Pro Leu Gln Leu Leu Lys Ser Met Gly Ala Pro Phe Ile

370 375 380

5 Phe Ile Lys Asn Trp Lys Glu Leu Pro Ala Val Leu Glu Lys Glu Lys

385 390 395 400

Thr Ile Ile Leu Gln Glu Lys Ile Glu Arg Arg Lys Met Leu Leu Gln

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35 40 45

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50 55 60

Glu Glu Thr Thr Phe Arg Met Glu Ser Asn Leu Tyr Gln Asp Gln Ser

65 70 75 80

Glu Asp Lys Arg Glu Lys Lys Glu Ala Lys Glu Lys Glu Lys Arg

5 85 90 95

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Asp Leu Glu Glu Lys Glu Pro Gly Asp His Glu Arg Ala Lys Ser Thr

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10 Val Met

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<212> DNA

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	gtcatcagca aaggctgcag ctgtaaactgc gtggatgact cacaggacta ctacgtggc	240
	aagaagaaca tcacgtgctg tgacaccgac ttgtgcacacg ccagcggggc ccatgcctg	300
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<212> DNA

5 <213> Homo sapiens

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<212> DNA

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25 atcagcaagg tgaatacagc agtccagtttta atcttggatgg cagttttttt ggcagcttca 780

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<213> Homo sapiens

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Gln Val Ser Asn Glu Asp Cys Leu Gln Val Lys Asn Cys Thr Gln Leu

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ggg gag cag tgc tgg acc ggc cgc atc cgc gca gtt ggc ctc ctg acc 195

Gly Glu Gln Cys Trp Thr Ala Arg Ile Arg Ala Val Gly Leu Leu Thr

c 15 45 50 55 60

gtc atc agc aaa ggc tgc agc ttg aac tgc gtg gat gac tca cag gac 243

Val Ile Ser Lys Gly Cys Ser Leu Asn Cys Val Asp Asp Ser Gln Asp

65 70 75

tac tac gtg ggc aag aag aac atc acg tgc tgt gac acc gac ttg tgc 291

20 Tyr Tyr Val Gly Lys Lys Asn Ile Thr Cys Cys Asp Thr Asp Leu Cys

80 85 90

aac gcc agc ggg gcc cat gcc ctg cag cgg gct gcc gcc atc ctt gcg 339

Asn Ala Ser Gly Ala His Ala Leu Gln Pro Ala Ala Ala Ile Leu Ala

95 100 105

25 ctg ctc cct gca ctc ggc ctg ctg ctc tgg gga ccc ggc cag cta 384

Leu Leu Pro Ala Leu Gly Leu Leu Leu Trp Gly Pro Gly Gin Leu

110 115 120

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5 cctccaaacc tctctgtgc tgggtccatg gcccaggcatt ctccacccctt aaccctgtgc	680
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15 <212> PRT

<213> Homo sapiens

<400> 22

Met Lys Ala Val Leu Leu Ala Leu Leu Met Ala Gly

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Leu Ala Leu Gln Pro Gly Thr Ala Leu Leu Cys Tyr Ser Cys Lys Ala

15	20	25
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Gln Val Ser Asn Glu Asp Cys Leu Gln Val Lys Asn Cys Thr Gln Leu

30	35	40
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25 Gly Glu Gln Cys Trp Thr Ala Arg Ile Arg Ala Val Gly Leu Leu Thr

45	50	55	60
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Val Ile Ser Lys Gly Cys Ser Leu Asn Cys Val Asp Asp Ser Gln Asp

65 70 75

Tyr Tyr Val Gly Lys Lys Asn Ile Thr Cys Cys Asp Thr Asp Leu Cys

80 85 90

Asn Ala Ser Gly Ala His Ala Leu Gln Pro Ala Ala Ala Ile Leu Ala

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Leu Leu Pro Ala Leu Gly Leu Leu Leu Trp Gly Pro Gly Gln Leu

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5 15 <400> 23

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ccgtccgtcc gtcggggcgc cgcagtcggccca gccaggccca gggccccgg cccctcgct 180

ccccgcaccc ggagccaccc ggtggagcgg gccttgccgc ggcagcc atg tcc atg 236

20 Met Ser Met

1

ggc ctg gag atc acg ggc acc gcg ctg gcc gtg ctg ggc tgg ctg ggc 284

Gly Leu Glu Ile Thr Gly Thr Ala Leu Ala Val Leu Gly Trp Leu Gly

5 10 15

25 acc atc gtg tgc tgc gcg ttg ccc atg tgg cgc gtg tgg gcc ttc atc 332

Thr Ile Val Cys Cys Ala Leu Pro Met Trp Arg Val Ser Ala Phe Ile

20 25 30 35

ggc agc aac atc atc acg tcg cag aac atc tgg gag ggc ctg tgg atg 380

Gly Ser Asn Ile Ile Thr Ser Gln Asn Ile Trp Glu Gly Leu Trp Met

40 45 50

aac tgc gtg gtg cag agc acc ggc cag atg cag tgc aag gtg tac gac 428

5 Asn Cys Val Val Gln Ser Thr Gly Gln Met Gln Cys Lys Val Tyr Asp

55 60 65

tcg ctg ctg gca ctg cca cag gac ctt cag gcg gcc cgc gcc ctc atc 476

Ser Leu Leu Ala Leu Pro Gln Asp Leu Gln Ala Ala Arg Ala Leu Ile

70 75 80

10 gtg gtg gcc atc ctg ctg gcc gcc ttc ggg ctg cta gtg gcg ctg gtg 524

Val Val Ala Ile Leu Leu Ala Ala Phe Gly Leu Leu Val Ala Leu Val

85 90 95

ggc gcc cag tgc acc aac tgc gtg cag gac gac acg gcc aag gcc aag 572

Gly Ala Gln Cys Thr Asn Cys Val Gln Asp Asp Thr Ala Lys Ala Lys

15 100 105 110 115

atc acc atc gtg gca ggc gtg ctg ttc ctt ctc gcc gcc ctg ctc acc 620

Ile Thr Ile Val Ala Gly Val Leu Phe Leu Leu Ala Ala Leu Leu Thr

120 125 130

ctc gtg ccg gtg tcc tgg tcg gcc aac acc att atc cgg gac ttc tac 668

20 Leu Val Pro Val Ser Trp Ser Ala Asn Thr Ile Ile Arg Asp Phe Tyr

135 140 145

aac ccc gtg gtg ccc gag ggc cag aag cgc gag atg ggc gcg ggc ctg 716

Asn Pro Val Val Pro Glu Ala Gln Lys Arg Glu Met Gly Ala Gly Leu

150 155 160

25 tac gtg ggc tgg gcg gcc ggc ctg cag ctg ctg ggg ggc gcg ctg 764

Tyr Val Gly Trp Ala Ala Ala Leu Gln Leu Leu Gly Gly Ala Leu

165 170 175

ctc tgc tgc tcg tgt ccc cca cgc gag aag aag tac acg gcc acc aag 812

Leu Cys Cys Ser Cys Pro Pro Arg Glu Lys Lys Tyr Thr Ala Thr Lys

180 185 190 195

gtc gtc tac tcc gcg ccg cgc tcc acc ggc ccg gga gcc agc ctg ggc 860

5 Val Val Tyr Ser Ala Pro Arg Ser Thr Gly Pro Gly Ala Ser Leu Gly

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aca ggc tac gac cgc aag gac tac gtc taa gggacagacg cagggagacc 910

Thr Gly Tyr Asp Arg Lys Asp Tyr Val

215 220

10 ccaccaccac caccaccacc aacaccacca ccaccacagc gagctggagc ggcacccagg 970

ccatccagcg tgcagccttg cctcgaggc cagccaccc ceagaagcca ggaagcccc 1030

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25 Met Ser Met

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Gly Ser Asn Ile Ile Thr Ser Gln Asn Ile Trp Glu Gly Leu Trp Met

5 40 45 50

Asn Cys Val Val Gln Ser Thr Gly Gln Met Gln Cys Lys Val Tyr Asp

55 60 65

Ser Leu Leu Ala-Leu Pro Gln Asp Leu Gln Ala Ala Arg Ala Leu Ile

70 75 80

10 15 20 25 30 35 40 45 50 55 60 65 70 75 80 85 90 95 100 105 110 115 120 125 130 135 140 145 150 155 160 165 170 175 180 185 190 195 200 205 210

Gly Ala Gln Cys Thr Asn Cys Val Gln Asp Asp Thr Ala Lys Ala Lys

Ile Thr Ile Val Ala Gly Val Leu Phe Leu Leu Ala Ala Leu Leu Thr

Leu Val Pro Val Ser Trp Ser Ala Asn Thr Ile Ile Arg Asp Phe Tyr

Asn Pro Val Val Pro Glu Ala Gln Lys Arg Glu Met Gly Ala Gly Leu

Tyr Val Gly Trp Ala Ala Ala Leu Gln Leu Leu Gly Gly Ala Leu

Leu Cys Cys Ser Cys Pro Pro Arg Glu Lys Lys Tyr Thr Ala Thr Lys

Val Val Tyr Ser Ala Pro Arg Ser Thr Gly Pro Gly Ala Ser Leu Gly

Thr Gly Tyr Asp Arg Lys Asp Tyr Val

215 220

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5 <212> DNA

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Met Ala Tyr His Gly Leu Thr Val Pro Leu Ile Val Met Ser Val

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Phe Trp Gly Phe Val Gly Phe Leu Val Pro Trp Phe Ile Pro Lys Gly

15 20 25 30

cct aac cgg gga gtt atc att acc atg ttg gtg acc tgt tca gtt tgc 203

Pro Asn Arg Gly Val Ile Ile Thr Met Leu Val Thr Cys Ser Val Cys

35 40 45

tgc tat ctc ttt tgg ctg att gca att ctg gcc caa ctc aac cct ctc 251

20 Cys Tyr Leu Phe Trp Leu Ile Ala Ile Leu Ala Gln Leu Asn Pro Leu

50 55 60

ttt gga ccg caa ttg aaa aat gaa acc atc tgg tat ctg aag tat cat 299

Phe Gly Pro Gln Leu Lys Asn Glu Thr Ile Trp Tyr Leu Lys Tyr His

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25 tgg cct tgagg aagaagacat gctctacagt gctcgttctt tgaggtcacg 350

Trp Pro

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cacgtgggaa	tcagtgaagt	gttttagaaac	tgcgtcaaga	caacaaagac	ccagtgggg	710	
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tatgg						835	

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15 <213> Homo sapiens

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20 25 30

Pro Asn Arg Gly Val Ile Ile Thr Met Leu Val Thr Cys Ser Val Cys

35 40 45

Cys Tyr Leu Phe Trp Leu Ile Ala Ile Leu Ala Gln Leu Asn Pro Leu

25 50 55 60

Phe Gly Pro Gln Leu Lys Asn Glu Thr Ile Trp Tyr Leu Lys Tyr His

65 70 75

Trp Pro

80

5 <210> 27

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ggcc atg cta gcc ttg cgc gtg gcg cgc ggc tcg tgg ggg gcc ctg cgc 169

Met Leu Ala Leu Arg Val Ala Arg Gly Ser Trp Gly Ala Leu Arg

15 1 5 10 15

ggc gcc gct tgg gct ccg gga acg cgg cgg agt aag cga cgc gcc tgc 217
Gly Ala Ala Trp Ala Pro Gly Thr Arg Pro Ser Lys Arg Arg Ala Cys

20 20 25 30

tgg gcc ctg ctg cgg ccc gtg ccc tgc tgc ttg ggc tgc ctg gcc gaa 265

20 Trp Ala Leu Leu Pro Pro Val Pro Cys Cys Leu Gly Cys Leu Ala Glu

35 40 45

cgc tgg agg ctg cgt ccg gcc gct ctt ggc ttg cgg ctg ccc ggg atc 313

Arg Trp Arg Leu Arg Pro Ala Ala Leu Gly Leu Arg Leu Pro Gly Ile

50 55 60

25 ggc cag cgg aac cac tgt tgc ggc gcg ggg aag ggc gct ccc agg cca 361

Gly Gln Arg Asn His Cys Ser Gly Ala Gly Lys Ala Ala Pro Arg Pro

65 70 75

gct gcc gga gct ggc gtc gct gcc gaa gcc ccc ggc ggc cag tgg ggc 409
Ala Ala Gly Ala Gly Ala Ala Ala Glu Ala Pro Gly Gly Gln Trp Gly
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ccg gct agc acc ccc agc ctg tat gaa aac cca tgg aca atc ccc aat 457
5 Pro Ala Ser Thr Pro Ser Leu Tyr Glu Asn Pro Trp Thr Ile Pro Asn
100 105 110

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Met Leu Ser Met Thr Arg Ile Gly Leu Ala Pro Val Leu Gly Tyr Leu
115 120 125

10 att att gaa gaa gat ttg aat att gca cta gga gtt ttg gct tta gct 553
Ile Ile Glu Glu Asp Phe Asn Ile Ala Leu Gly Val Phe Ala Leu Ala
130 135 140

gga cta aca gat ttg ttg gat gga ttg att gct cga aac tgg gcc aat 601
Gly Leu Thr Asp Leu Leu Asp Gly Phe Ile Ala Arg Asn Trp Ala Asn
145 150 155

15 caa aga tca gct ttg gga agt gct ctt gat cca ctt gct gat aaa ata 649
Gln Arg Ser Ala Leu Gly Ser Ala Leu Asp Pro Leu Ala Asp Lys Ile
160 165 170 175

ctt atc agt atc tta tat gtt agc ttg acc tat gca gat ctt att cca 697
20 Leu Ile Ser Ile Leu Tyr Val Ser Leu Thr Tyr Ala Asp Leu Ile Pro
180 185 190

gtt cca ctt act tac atg atc att tgg aga gat gta atg ttg att gct 745
Val Pro Leu Thr Tyr Met Ile Ile Ser Arg Asp Val Met Leu Ile Ala
195 200 205

25 gct gtt ttg tat gtc aga tac cga act ctt cca aca cca cga aca ctt 793
Ala Val Phe Tyr Val Arg Tyr Arg Thr Leu Pro Thr Pro Arg Thr Leu
210 215 220

gcc aag tat ttc aat cct tgc tat gcc act gct agg tta aaa cca aca 841

Ala Lys Tyr Phe Asn Pro Cys Tyr Ala Thr Ala Arg Leu Lys Pro Thr

225 230 235

ttc atc agc aag gtg aat aca gca gtc cag tta atc ttg gtg gca gct 889

5 Phe Ile Ser Lys Val Asn Thr Ala Val Gln Leu Ile Leu Val Ala Ala

240 245 250 255

tct ttg gca gct cca gtt ttc aac tat gct gac agc att tat ctt cag 937

Ser Leu Ala Ala Pro Val Phe Asn Tyr Ala Asp Ser Ile Tyr Leu Gln

260 265 270

10 ata cta tgg tgt ttt aca gct ttc acc aca gct gca tca gct tat agt 985

Ile Leu Trp Cys Phe Thr Ala Phe Thr Thr Ala Ala Ser Ala Tyr Ser

275 280 285

tac tat cat tat ggc cgg aag act gtt cag gtg ata aaa gac tga 1030

Tyr Tyr His Tyr Gly Arg Lys Thr Val Gln Val Ile Lys Asp

15 290 295 300

tgaaaatgtcat ccctcaactgt tagtaaggaa gcagtataaca tcaatggaa cagggccat 1090

ggaaatgtac aggagtttcc ctattttgtt gttagcttg aaaaaggact tgcagaatc 1150

aactgtgtca tcaaaattta agtaatgtgc attgaaaata aggttgatca tggaaatatg 1210

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25 <213> Homo sapiens

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20

25

30

5 Trp Ala Leu Leu Pro Pro Val Pro Cys Cys Leu Gly Cys Leu Ala Glu

35

40

45

Arg Trp Arg Leu Arg Pro Ala Ala Leu Gly Leu Arg Leu Pro Gly Ile

50

55

60

Gly Gln Arg Asn His Cys Ser Gly Ala Gly Lys Ala Ala Pro Arg Pro

10

65

70

75

Ala Ala Gly Ala Gly Ala Ala Glu Ala Pro Gly Gly Gln Trp Gly

80 85 90 95

Pro Ala Ser Thr Pro Ser Leu Tyr Glu Asn Pro Trp Thr Ile Pro Asn

100

105

110

15 Met Leu Ser Met Thr Arg Ile Glv Leu Ala Pro Val Leu Gly Tyr Leu

115

120

125

Ile Ile Glu Glu Asp Phe Asn Ile Ala Leu Gly Val Phe Ala Leu Ala

130

135

140

Gly Leu Thr Asp Leu Leu Asp Gly Phe Ile Ala Arg Asn Trp Ala Asn

20

145

150

155

Gln Arg Ser Ala Leu Gly Ser Ala Leu Asp Pro Leu Ala Asp Lys Ile

160 165 170 175

Leu Ile Ser Ile Leu Tyr Val Ser Leu Thr Tyr Ala Asp Leu Ile Pro

180

185

190

25 Val Pro Leu Thr Tyr Met Ile Ile Ser Arg Asp Val Met Leu Ile Ala

195

200

205

Ala Val Phe Tyr Val Arg Tyr Arg Thr Leu Pro Thr Pro Arg Thr Leu

210 215 220

Ala Lys Tyr Phe Asn Pro Cys Tyr Ala Thr Ala Arg Leu Lys Pro Thr

225 230 235

Phe Ile Ser Lys Val Asn Thr Ala Val Gln Leu Ile Leu Val Ala Ala

5 240 245 250 255

Ser Leu Ala Ala Pro Val Phe Asn Tyr Ala Asp Ser Ile Tyr Leu Gln

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275 280 285

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15 <211> 1305

<212> DNA

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<400> 29

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1 5 10

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Asp Ala Tyr Pro Lys Thr Leu Glu Asp Phe Arg Val Lys Thr Cys Gly

25 15 20 25

ggc gcc acc gtg acc att gtc agt ggc ctt ctc atg ctg cta ctg ttc 147

Gly Ala Thr Val Thr Ile Val Ser Gly Leu Leu Met Leu Leu Phe

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	ctg tcc gag ctg cag tat tac ctc acc acg gag gtg cat cct gag ctc			195
	Leu Ser Glu Leu Gln Tyr Tyr Leu Thr Thr Glu Val His Pro Glu Leu			---
	45	50	55	
5	tac gtg gac aag tcg cgg gga gat aaa ctg aag atc aac atc gat gta			243
	Tyr Val Asp Lys Ser Arg Gly Asp Lys Leu Lys Ile Asn Ile Asp Val			
	60	65	70	
	ctt ttt ccg cac atg cct tgt gcc tat ctg agt att gat gcc atg gat			291
	Leu Phe Pro His Met Pro Cys Ala Tyr Leu Ser Ile Asp Ala Met Asp			
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	gtg gcc gga gaa cag cag ctg gat gtg gaa cac aac ctg ttc aag caa			339
	Val Ala Gly Glu Gln Gln Leu Asp Val Glu His Asn Leu Phe Lys Gln			
	95	100	105	
	cga cta gat aaa gat ggc atc ccc gtg agc tca gag gct gag cgg cat			387
15	Arg Leu Asp Lys Asp Gly Ile Pro Val Ser Ser Glu Ala Glu Arg His			
	110	115	120	
	gag ctt ggg aaa gtc gag gtg acg gtg ttt gac cct gac tcc ctg gac			435
	Glu Leu Gly Lys Val Glu Val Thr Val Phe Asp Pro Asp Ser Leu Asp			
	125	130	135	
20	cct gat cgc tgt gag agc tgc tat ggt gct gag gca gaa gat atc aag			483
	Pro Asp Arg Cys Glu Ser Cys Tyr Gly Ala Glu Ala Glu Asp Ile Lys			
	140	145	150	
	tgc tgt aac acc tgt gaa gat gtg cgg gag gca tat cgc cgt aga ggc			531
	Cys Cys Asn Thr Cys Glu Asp Val Arg Glu Ala Tyr Arg Arg Gly			
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	tgg gcc ttc aag aac cca gat act att gag cag tgc cgg cga gag ggc			579
	Trp Ala Phe Lys Asn Pro Asp Thr Ile Glu Gln Cys Arg Arg Glu Gly			

	175	180	185	
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	Phe Ser Gln Lys Met Gln Glu Gln Lys Asn Glu Gly Cys Gln Val Tyr			
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	Gly Phe Leu Glu Val Asn Lys Val Ala Gly Asn Phe His Phe Ala Pro			
	205	210	215	
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	Gly Lys Ser Phe Gln Gln Ser His Val His Val His Asp Leu Gln Ser			
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	ttt ggc ctt gac aac atc aac atg acc cac tac atc cag cac ctg tca		771	
	Phe Gly Leu Asp Asn Ile Asn Met Thr His Tyr Ile Gln His Leu Ser			
	235	240	245	250
	ttt ggg gag gac tat cca ggc att gtg aac ccc ctg gac cac acc aat		819	
15	Phe Gly Glu Asp Tyr Pro Gly Ile Val Asn Pro Leu Asp His Thr Asn			
	255	260	265	
	gtc act ggc ccc caa gcc tcc atg atg ttc cag tac ttt gtg aag gtg		867	
	Val Thr Ala Pro Gln Ala Ser Met Met Phe Gln Tvr Phe Val Lys Val			
	270	275	280	
20	gtg ccc act gtg tac atg aag gtg gac gga gag gta ctg agg aca aat		915	
	Val Pro Thr Val Tyr Met Lys Val Asp Gly Glu Val Leu Arg Thr Asn			
	285	290	295	
	cag ttc tct gtg acc aga cat gag aag gtt gcc aat ggg ctg ttg ggc		963	
	Gln Phe Ser Val Thr Arg His Glu Lys Val Ala Asn Gly Leu Leu Gly			
25	300	305	310	
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315 320 325 330
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Asp Ser Leu Ile Tyr His Ser Ala Arg Ala Ile Gln Lys Lys Ile Asp
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cta ggg aag aca acg tagtccacct cggtgcttcc tctgtctctt ctttctccct 1210
Leu Gly Lys Thr Thr
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Leu Ser Glu Leu Gln Tyr Tyr Leu Thr Thr Glu Val His Pro Glu Leu

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5 Tyr Val Asp Lys Ser Arg Gly Asp Lys Leu Lys Ile Asn Ile Asp Val

60 65 70

Leu Phe Pro His Met Pro Cys Ala Tyr Leu Ser Ile Asp Ala Met Asp

75 80 85 90

Val Ala Gly Glu Gln Gln Leu Asp Val Glu His Asn Leu Phe Lys Gln

10 95 100 105

Arg Leu Asp Lys Asp Gly Ile Pro Val Ser Ser Glu Ala Glu Arg His

110 115 120

Glu Leu Gly Lys Val Glu Val Thr Val Phe Asp Pro Asp Ser Leu Asp

125 130 135

15 Pro Asp Arg Cys Glu Ser Cys Tyr Gly Ala Glu Ala Glu Asp Ile Lys

140 145 150

Cys Cys Asn Thr Cys Glu Asp Val Arg Glu Ala Tyr Arg Arg Arg Gly

155 160 165 170

Trp Ala Phe Lys Asn Pro Asp Thr Ile Glu Gln Cys Arg Arg Glu Gly

20 175 180 185

Phe Ser Gln Lys Met Gln Glu Gln Lys Asn Glu Gly Cys Gln Val Tyr

190 195 200

Gly Phe Leu Glu Val Asn Lys Val Ala Gly Asn Phe His Phe Ala Pro

205 210 215

25 Gly Lys Ser Phe Gln Gln Ser His Val His Val His Asp Leu Gln Ser

220 225 230

Phe Gly Leu Asp Asn Ile Asn Met Thr His Tyr Ile Gln His Leu Ser

235 240 245 250

Phe Gly Glu Asp Tyr Pro Gly Ile Val Asn Pro Leu Asp His Thr Asn

255 260 265

Val Thr Ala Pro Gln Ala Ser Met Met Phe Gln Tyr Phe Val Lys Val

5 270 275 280

Val Pro Thr Val Tyr Met Lys Val Asp Gly Glu Val Leu Arg Thr Asn

285 290 295

Gln Phe Ser Val Thr Arg His Glu Lys Val Ala Asn Gly Leu Leu Gly

300 305 310

6 10 Asp Gln Gly Leu Pro Gly Val Phe Val Leu Tyr Glu Leu Ser Pro Met

315 320 325 330

Met Val Lys Leu Thr Glu Lys His Arg Ser Phe Thr His Phe Leu Thr

335 340 345

Gly Val Cys Ala Ile Ile Gly Gly Met Phe Thr Val Ala Gly Leu Ile

6 15 350 355 360

Asp Ser Leu Ile Tyr His Ser Ala Arg Ala Ile Gln Lys Lys Ile Asp

365 370 375

Leu Gly Lys Thr Thr

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20

<210> 31

<211> 899

<212> DNA

25 <213> Homo sapiens

<400> 31

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 agacttctgc gtt atg acc cgg ctg ctg ggc tac gtg gac ccc ctg gat 109

Met Thr Arg Leu Leu Gly Tyr Val Asp Pro Leu Asp

1 5 10

5 ccc agc ttt gtg gct gcc gtc atc acc atc acc ttc aat ccg ctc tac 157

Pro Ser Phe Val Ala Ala Val Ile Thr Ile Thr Phe Asn Pro Leu Tyr

15 20 25

tgg aat gtg gtt gca cga tgg gaa cac aag acc cgc aag ctg agc agg 205

Trp Asn Val Val Ala Arg Trp Glu His Lys Thr Arg Lys Leu Ser Arg

10 30 35 40

gcc ttc gga tcc ccc tac ctg gcc tgc tac tct cta agc gtc acc atc 253

Ala Phe Gly Ser Pro Tyr Leu Ala Cys Tyr Ser Leu Ser Val Thr Ile

45 50 55 60

ctg ctc ctg aac ttc ctg cgc tcg cac tgc ttc acg cag gag gcc atg ctg 301

15 Leu Leu Leu Asn Phe Leu Arg Ser His Cys Phe Thr Gln Ala Met Leu

65 70 75

agc cag ccc agg atg gag agc ctg gac acc ccc gcg gcc tac agc ctg 349

Ser Gln Pro Arg Met Glu Ser Leu Asp Thr Pro Ala Ala Tyr Ser Leu

80 85 90

20 ggc ctc gcg ctc ctg gga ctg ggc gtc gtg ctc gtg ctc tcc agc ttc 397

Gly Leu Ala Leu Leu Gly Leu Gly Val Val Leu Val Leu Ser Ser Phe

95 100 105

ttt gca ctg ggg ttc gct gga act ttc cta ggt gat tac ttc ggg atc 445

Phe Ala Leu Gly Phe Ala Gly Thr Phe Leu Gly Asp Tyr Phe Gly Ile

25 110 115 120

ctc aag gag gcg aga gtg acc gtg ttc ccc ttc aac atc ctg gac aac 493

Leu Lys Glu Ala Arg Val Thr Val Phe Pro Phe Asn Ile Leu Asp Asn

125

130

135

140

ccc atg tac tgg gga agc aca gcc aac tac ctg ggc tgg gcc atc atg 541

Pro Met Tyr Trp Gly Ser Thr Ala Asn Tyr Leu Gly Trp Ala Ile Met

145

150

155

5 cac gcc agc ccc acg ggc ctg ctc ctg acg gtg ctg gtg gcc ctc acc 589

His Ala Ser Pro Thr Gly Leu Leu Leu Thr Val Leu Val Ala Leu Thr

160

165

170

tac ata gtg gct ctc cta tac gaa gag ccc ttc acc gct gag atc tac 637

Tyr Ile Val Ala Leu Leu Tyr Glu Glu Pro Phe Thr Ala Glu Ile Tyr

10

175

180

185

cg~~g~~ cag aaa gcc tcc ggg tcc cac aag agg agc tgattgagct gcaacagct 690

Arg Gln Lys Ala Ser Gly Ser His Lys Arg Ser

190

195

tgctgaaggc ctggccagcc tccctggccctg ccccaagtgg caggccctgc gcagggcgcag 750

15

aatggtgccct gctgctcagg gctcgcccc ggcgtgggct gccccagtgc cttggaaacct 810

gctgccttgg ggaccttggg cgtgcccaca tatggccatt gagtccaaac ccacacattc 870

ccattcacca ataaaggcac cctgaccccc 899

20 <210> 32

<211> 199

<212> PRT

<213> Homo sapiens

25 <400> 32

Met Thr Arg Leu Leu Gly Tyr Val Asp Pro Leu Asp

1

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10

Pro Ser Phe Val Ala Ala Val Ile Thr Ile Thr Phe Asn Pro Leu Tyr

15 20 25

Trp Asn Val Val Ala Arg Trp Glu His Lys Thr Arg Lys Leu Ser Arg

30 35 40

5 Ala Phe Gly Ser Pro Tyr Leu Ala Cys Tyr Ser Leu Ser Val Thr Ile

45 50 55 60

Leu Leu Leu Asn Phe Leu Arg Ser His Cys Phe Thr Gln Ala Met Leu

65 70 75

Ser Gln Pro Arg Met Glu Ser Leu Asp Thr Pro Ala Ala Tyr Ser Leu

10 80 85 90

Gly Leu Ala Leu Leu Gly Leu Gly Val Val Leu Val Leu Ser Ser Phe

95 100 105

Phe Ala Leu Gly Phe Ala Gly Thr Phe Leu Gly Asp Tyr Phe Gly Ile

110 115 120

15 125 130 135 140

Pro Met Tyr Trp Gly Ser Thr Ala Asn Tyr Leu Gly Trp Ala Ile Met

145 150 155

His Ala Ser Pro Thr Gly Leu Leu Leu Thr Val Leu Val Ala Leu Thr

20 160 165 170

Tyr Ile Val Ala Leu Leu Tyr Glu Glu Pro Phe Thr Ala Glu Ile Tyr

175 180 185

Arg Gln Lys Ala Ser Gly Ser His Lys Arg Ser

190 195

25

<211> 905

<212> DNA

<213> Homo sapiens

5 <400> 33

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Met Ala Pro His Gly Pro Glv Ser Leu Thr Thr Leu Val Pro

1 - 5 10

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10 Trp Ala Ala Ala Leu Leu Leu Ala Leu Glv Val Glu Arg Ala Leu Ala

15 20 25 30

cta ccc gag ata tgc acc caa tgt cca ggg agc gtg caa aat ttg tca 146

Leu Pro Glu Ile Cys Thr Gln Cys Pro Gly Ser Val Gln Asn Leu Ser

35 40 45

15 aaa gtg gcc ttt tat tgt aaa acg aca cga gag cta atg ctg cat gcc 194

Lys Val Ala Phe Tyr Cys Lys Thr Thr Arg Glu Leu Met Leu His Ala

50 55 60

cgt tgc tgc ctg aat cag aag ggc acc atc ttg ggg ctg gat ctc cag 242

Arg Cys Cys Leu Asn Gln Lys Gly Thr Ile Leu Gly Leu Asp Leu Gln

20 65 70 75

aac tgt tct ctg gag gac cct ggt cca aac ttt cat cag gca cat acc 290

Asn Cys Ser Leu Glu Asp Pro Gly Pro Asn Phe His Gln Ala His Thr

80 85 90

act gtc atc ata gac ctg caa gca aac ccc ctc aaa ggt gac ttg gcc 338

25 Thr Val Ile Ile Asp Leu Gln Ala Asn Pro Leu Lys Gly Asp Leu Ala

95 100 105 110

aac acc ttc cgt ggc ttt act cag ctc cag act ctg ata ctg cca caa 386

Asn Thr Phe Arg Gly Phe Thr Gln Leu Gln Thr Leu Ile Leu Pro Gln

115 120 125

cat gtc aac tgt cct gga gga att aat gcc tgg aat act atc acc tct 434

His Val Asn Cys Pro Gly Gly Ile Asn Ala Trp Asn Thr Ile Thr Ser

5 130 135 140

tat ata gac aac caa atc tgt caa ggg caa aag aac ctt tgc aat aac 482

Tyr Ile Asp Asn Gln Ile Cys Gln Gly Gln Lys Asn Leu Cys Asn Asn

145 150 155

act ggg gac cca gaa atg tgt cct gag aat gga tct tgt gta cct gat 530

10 Thr Glv Asp Pro Glu Met Cys Pro Glu Asn Glv Ser Cys Val Pro Asp

160 165 170

ggt cca ggt ctt ttg cag tgt gtt tgt gct gat ggt ttc cat gga tac 578

Gly Pro Gly Leu Leu Gln Cys Val Cys Ala Asp Gly Phe His Gly Tyr

175 180 185 190

15 aag tgt atg cgc cag ggc tcg ttc tca ctg ctt atg ttc ttc ggg att 626

Lys Cys Met Arg Gln Gly Ser Phe Ser Leu Leu Met Phe Phe Gly Ile

195 200 205

ctg gga gcc acc act cta tcc gtc tcc att ctg ctt tgg gcg acc cag 674

Leu Gly Ala Thr Thr Leu Ser Val Ser Ile Leu Leu Trp Ala Thr Gln

20 210 215 220

cgc cga aaa gcc aag act tca tgaac tacataggct ttaccattga 720

Arg Arg Lys Ala Lys Thr Ser

225

cctaagatca atctgaacta tcttagccca gtcaggagc tctgcttctt agaaaaggcat 780

25 ctttcgccag tggattgcc tcaagggttga gccccccatt ggaagatgaa aaattgcact 840

cccttggtgt agacaaatac cagttcccat tggtgttgcctataata aacacttttt 900

ctttt 905

<210> 34

<211> 229

5 <212> PRT

<213> Homo sapiens

<400> 34

Met Ala Pro His Gly Pro Gly Ser Leu Thr Thr Leu Val Pro

6310	1	5	10
6311	Trp Ala Ala Ala Leu Leu Leu Ala Leu Gly Val Glu Arg Ala Leu Ala		
6312	15	20	25
6313	Leu Pro Glu Ile Cys Thr Gln Cys Pro Gly Ser Val Gln Asn Leu Ser		30
6314	35	40	45
6315	Lys Val Ala Phe Tyr Cys Lys Thr Thr Arg Glu Leu Met Leu His Ala		
6316	50	55	60
6317	Arg Cys Cys Leu Asn Gln Lys Gly Thr Ile Leu Gly Leu Asp Leu Gln		
6318	65	70	75
6319	Asn Cys Ser Leu Glu Asp Pro Gly Pro Asn Phe His Gln Ala His Thr		
20	80	85	90
6320	Thr Val Ile Ile Asp Leu Gln Ala Asn Pro Leu Lys Gly Asp Leu Ala		
6321	95	100	105
6322	Asn Thr Phe Arg Gly Phe Thr Gln Leu Gln Thr Leu Ile Leu Pro Gln		110
6323	115	120	125
25	His Val Asn Cys Pro Gly Gly Ile Asn Ala Trp Asn Thr Ile Thr Ser		
6324	130	135	140
6325	Tyr Ile Asp Asn Gln Ile Cys Gln Gly Gln Lys Asn Leu Cys Asn Asn		

145 150 155

Thr Gly Asp Pro Glu Met Cys Pro Glu Asn Gly Ser Cys Val Pro Asp

160 165 170

Gly Pro Gly Leu Leu Gln Cys Val Cys Ala Asp Gly Phe His Gly Tyr

5 175 180 185 190

Lys Cys Met Arg Gln Gly Ser Phe Ser Leu Leu Met Phe Phe Gly Ile

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Leu Gly Ala Thr Thr Leu Ser Val Ser Ile Leu Leu Trp Ala Thr Gln

210 215 220

6 10 Arg Arg Lys Ala Lys Thr Ser

225

<210> 35

15 <211> 841

<212> DNA

<213> Homo sapiens

<400> 35

20 ctccacgagg ctgcgggtt aggacccca gctccgac atg tct ccc tct ggt cgc 56

Met Ser Pro Ser Gly Arg

1 5

ctg tgt ctt ctc acc atc gtt ggc ctg att ctc ccc acc aga gga cag 104

Leu Cys Leu Leu Thr Ile Val Gly Leu Ile Leu Pro Thr Arg Gly Gin

25 10 15 20

acg ttg aaa gat acc acg tcc agt tct tca gca gac tca act atc atg 152

Thr Leu Lys Asp Thr Thr Ser Ser Ser Ala Asp Ser Thr Ile Met

	25	30	35	
	gac att cag gtc ccg aca cga gcc cca gat gca gtc tac aca gaa ctc			200
	Asp Ile Gln Val Pro Thr Arg Ala Pro Asp Ala Val Tyr Thr Glu Leu			—
	40	45	50	
5	cag ccc acc tct cca acc cca acc tgg cct gct gat gaa aca cca caa			248
	Gln Pro Thr Ser Pro Thr Pro Trp Pro Ala Asp Glu Thr Pro Gln			
	55	60	65	70
	ccc cag acc cag acc cag caa ctg gaa gga acg gat ggg cct cta gtg			296
	Pro Gln Thr Gln Thr Gln Gln Leu Glu Gly Thr Asp Gly Pro Leu Val			
10		75	80	85
	aca gat cca gag aca cac aag agc acc aaa gca gct cat ccc act gat			344
	Thr Asp Pro Glu Thr His Lys Ser Thr Lys Ala Ala His Pro Thr Asp			
	90	95	100	
	gac acc acg acg ctc tct gag aga cca tcc cca agc aca gac gtc cag			392
15	Asp Thr Thr Thr Leu Ser Glu Arg Pro Ser Pro Ser Thr Asp Val Gln			
	105	110	115	
	aca gac ccc cag acc ctc aag cca tct ggt ttt cat gag gat gac ccc			440
	Thr Asp Pro Gln Thr Leu Lys Pro Ser Gly Phe His Glu Asp Asp Pro			
	120	125	130	
20	ttc ttc tat gat gaa cac acc ctc egg aaa cgg ggg ctg ttg gtc gca			488
	Phe Phe Tyr Asp Glu His Thr Leu Arg Lys Arg Gly Leu Leu Val Ala			
	135	140	145	150
	gct gtg ctg ttc atc aca ggc atc atc atc ctc acc agt ggc aag tgc			536
	Ala Val Leu Phe Ile Thr Gly Ile Ile Ile Leu Thr Ser Gly Lys Cys			
25		155	160	165
	agg cag ctg tcc cgg tta tgc cgg aat cat tgc agg tgagtcca			580
	Arg Gln Leu Ser Arg Leu Cys Arg Asn His Cys Arg			

170

175

tcagaaacag gagctgacaa cccgctggc acccgaaagac caagccccct gccagctcac	640
cgtgcccagc ctccctgcata ccctcgaaga gcctggccag agagggaaaga cacagatgat	700
gaagctggag ccagggctgc cggtccgagt ctccctacctc ccccaacctt gcccggccct	760
5 gaaggctacc tggcgcccttg ggggctgtcc ctcaagttat ctccctgtt aagacaaaaa	820
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<210> 36

10 <211> 178

<212> PRT

<213> Homo sapiens

<400> 36

Met Ser Pro Ser Gly Arg

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Leu Cys Leu Leu Thr Ile Val Gly Leu Ile Leu Pro Thr Arg Gly Gln

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20

Thr Leu Lys Asp Thr Thr Ser Ser Ser Ala Asp Ser Thr Ile Met

20 25

30

35

Asp Ile Gln Val Pro Thr Arg Ala Pro Asp Ala Val Tyr Thr Glu Leu

40

45

50

Gln Pro Thr Ser Pro Thr Pro Trp Pro Ala Asp Glu Thr Pro Gln

55

60

65

70

25 Pro Gln Thr Gln Thr Gln Gln Leu Glu Gly Thr Asp Gly Pro Leu Val

75

80

85

Thr Asp Pro Glu Thr His Lys Ser Thr Lys Ala Ala His Pro Thr Asp

90

95

100

Asp Thr Thr Thr Leu Ser Glu Arg Pro Ser Pro Ser Thr Asp Val Gln

105

110

115

Thr Asp Pro Gln Thr Leu Lys Pro Ser Gly Phe His Glu Asp Asp Pro

5

120

125

130

Phe Phe Tyr Asp Glu His Thr Leu Arg Lys Arg Gly Leu Leu Val Ala

135

140

145

150

Ala Val Leu Phe Ile Thr Gly Ile Ile Ile Leu Thr Ser Gly Lys Cys

155

160

165

10 Arg Gln Leu Ser Arg Leu Cys Arg Asn His Cys Arg

170

175

<210> 37

<211> 1451

<212> DNA

<213> Homo sapiens

<400> 37

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gccccgttcca tggcggttggc tgccctgactg gaagcccgag tggg atg cgg ctg acg 116
Met Arg Leu Thr

1

cgg aag cgg ctc tgc tcg ttt ctt atc gcc ctg tac tgc cta ttc tcc 164
25 Arg Lys Arg Leu Cys Ser Phe Leu Ile Ala Leu Tyr Cys Leu Phe Ser
5 10 15 20
ctc tac gct gcc tac cac gtc ttc ttc ggg cgc cgc cgc cag gcg cgc 212

Leu Tyr Ala Ala Tyr His Val Phe Phe Gly Arg Arg Arg Gln Ala Pro

25 30 35

gcc ggg tcc ccg cgg ggc ctc agg aag ggg gcg gcc ccc gcg cgg gag 260

Ala Gly Ser Pro Arg Gly Leu Arg Lys Gly Ala Ala Pro Ala Arg Glu

5 40 45 50

aga cgc ggc cga gaa cag tcc act ttg gaa agt gaa gaa tgg aat cct 308

Arg Arg Gly Arg Glu Gln Ser Thr Leu Glu Ser Glu Glu Trp Asn Pro

55 60 65

tgg gaa gga gat gaa aaa aat gag caa caa cac aga ttt aaa act agc 356

10 Trp Glu Gly Asp Glu Lys Asn Glu Gln Gln His Arg Phe Lys Thr Ser

70 75 80

ctt caa ata tta gat aaa tcc acg aaa gga aaa aca gat ctc agt gta 404

Leu Gln Ile Leu Asp Lys Ser Thr Lys Gly Lys Thr Asp Leu Ser Val

85 90 95 100

caa atc tgg ggc aaa gct gcc att ggc ttt tat ctc tgg gag cat att 452

Gln Ile Trp Gly Lys Ala Ala Ile Gly Leu Tyr Leu Trp Glu His Ile

105 110 115

ttt gaa ggc tra ctt gat ccc agc gat gtg act gct caa tgg aga gaa 500

Phe Glu Gly Leu Leu Asp Pro Ser Asp Val Thr Ala Gln Trp Arg Glu

20 120 125 130

gga aag tca atc gta gga aga aca cag tac agc ttc atc act ggt cca 548

Gly Lys Ser Ile Val Gly Arg Thr Gln Tyr Ser Phe Ile Thr Gly Pro

135 140 145

gct gta ata cca ggg tac ttc tcc gtt gat gtg aat aat gtg gta ctc 596

25 Ala Val Ile Pro Gly Tyr Phe Ser Val Asp Val Asn Asn Val Val Leu

150 155 160

att tta aat gga aga gaa aaa gca aag atc ttt tat gcc acc cag tgg 644

ile leu asn gly arg glu lys ala lys ile phe tyr ala thr gln trp
 165 170 175 180
 tta ctt tat gca caa aat tta gtg caa att caa aaa ctc cag cat ctt 692
 Leu Leu Tyr Ala Gln Asn Leu Val Gln Ile Gln Lys Leu Gln His Leu
 5 185 190 195
 gct gtt gtt ttg ctc gga aat gaa cat tgt gat aat gag tgg ata aac 740
 Ala Val Val Leu Leu Gly Asn Glu His Cys Asp Asn Glu Trp Ile Asn
 200 - 205 210
 cca ttc ctc aaa aga aat gga ggc ttc gtg gag ctg ctt ttc ata ata 788
 10 Pro Phe Leu Lys Arg Asn Gly Gly Phe Val Glu Leu Leu Phe Ile Ile
 215 220 225
 tat gac agc ccc tgg att aat gac gtg gat gtt ttt cag tgg cct tta 836
 Tyr Asp Ser Pro Trp Ile Asn Asp Val Asp Val Phe Gln Trp Pro Leu
 230 235 240
 15 gga gta gca aca tac agg aat ttt cct gtg gtg gag gca agt tgg tca 884
 Gly Val Ala Thr Tyr Arg Asn Phe Pro Val Val Glu Ala Ser Trp Ser
 245 250 255 260
 atg ctg cat gat gag agg cca tat tta tgt aat ttc tta gga acg att 932
 Met Leu His Asp Glu Arg Pro Tyr Leu Cys Asn Phe Leu Gly Thr Ile
 20 265 270 275
 tat gaa aat tca tcc aga cag gca cta atg aac att ttg aaa aaa gat 980
 Tyr Glu Asn Ser Ser Arg Gln Ala Leu Met Asn Ile Leu Lys Lys Asp
 280 285 290
 ggg aac gat aag ctt tgt tgg gtt tca gca aga gaa cac tgg cag cct 1028
 25 Gly Asn Asp Lys Leu Cys Trp Val Ser Ala Arg Glu His Trp Gln Pro
 295 300 305
 cag gaa aca aat gaa agt ctt aag aat tac caa gat gcc ttg ctt cag 1076

Gln Glu Thr Asn Glu Ser Leu Lys Asn Tyr Gln Asp Ala Leu Leu Gln

310

315

320

agt gat ctc aca ttg tgc ccg gtc gga gta aac aca gaa tgc tat cga 1124

Ser Asp Leu Thr Leu Cys Pro Val Gly Val Asn Thr Glu Cys Tyr Arg

5 325

330

335

340

atc tat gag gct tgc tcc tat ggc tcc att cct gtg gtg gaa gac gtg 1172

Ile Tyr Glu Ala Cys Ser Tyr Gly Ser Ile Pro Val Val Glu Asp Val

345

350

355

atg aca gct ggc aac tgt ggg aat aca tct gtg cac cac ggt gct cct 1220

10 Met Thr Ala Gly Asn Cys Gly Asn Thr Ser Val His His Gly Ala Pro

360

365

370

ctg cag tta ctc aag tcc atg ggt gct ccc ttt atc ttt atc aag aac 1268

Leu Gln Leu Leu Lys Ser Met Gly Ala Pro Phe Ile Phe Ile Lys Asn

375

380

385

tgg aag gaa ctc cct gct gtt tta gaa aaa gag aaa act ata att tta 1316

Trp Lys Glu Leu Pro Ala Val Leu Glu Lys Glu Lys Thr Ile Ile Leu

390

395

400

caa gaa aaa att gaa aga aga aaa atg tta ctt cag tgg tat cag cac 1364

Gln Glu Lys Ile Glu Arg Arg Lys Met Leu Leu Gln Trp Tyr Gln His

20 405

410

415

420

ttc aag aca gag ctt aaa atg aaa ttt act aat att tta gaa agc tca 1412

Phe Lys Thr Glu Leu Lys Met Lys Phe Thr Asn Ile Leu Glu Ser Ser

425

430

435

ttt tta atg aat aat aaa agt taattat ctttttgagc t

1451

25 Phe Leu Met Asn Asn Lys Ser

440

<210> 38

<211> 443

<212> PRT

5 <213> Homo sapiens

<400> 38

Met Arg Leu Thr

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10 Arg Lys Arg Leu Cys Ser Phe Leu Ile Ala Leu Tyr Cys Leu Phe Ser

5 10 15 20

Leu Tyr Ala Ala Tyr His Val Phe Phe Gly Arg Arg Gln Ala Pro

25 30 35

Ala Gly Ser Pro Arg Gly Leu Arg Lys Gly Ala Ala Pro Ala Arg Glu

15 40 45 50

Arg Arg Gly Arg Glu Gln Ser Thr Leu Glu Ser Glu Glu Trp Asn Pro

55 60 65

Trp Glu Gly Asp Glu Lys Asn Glu Gln Gln His Arg Phe Lys Thr Ser

70 75 80

20 Leu Gln Ile Leu Asp Lys Ser Thr Lys Gly Lys Thr Asp Leu Ser Val

85 90 95 100

Gln Ile Trp Gly Lys Ala Ala Ile Gly Leu Tyr Leu Trp Glu His Ile

105 110 115

Phe Glu Gly Leu Leu Asp Pro Ser Asp Val Thr Ala Gln Trp Arg Glu

25 120 125 130

Gly Lys Ser Ile Val Gly Arg Thr Gln Tyr Ser Phe Ile Thr Gly Pro

135 140 145

Ala Val Ile Pro Gly Tyr Phe Ser Val Asp Val Asn Asn Val Val Leu

150 155 160

Ile Leu Asn Gly Arg Glu Lys Ala Lys Ile Phe Tyr Ala Thr Gln Trp

165 170 175 180

5 Leu Leu Tyr Ala Gln Asn Leu Val Gln Ile Gln Lys Leu Gln His Leu

185 190 195

Ala Val Val Leu Leu Gly Asn Glu His Cys Asp Asn Glu Trp Ile Asn

200 205 210

Pro Phe Leu Lys Arg Asn Gly Gly Phe Val Glu Leu Leu Phe Ile Ile

10 215 220 225

Tyr Asp Ser Pro Trp Ile Asn Asp Val Asp Val Phe Gln Trp Pro Leu

230 235 240

Gly Val Ala Thr Tyr Arg Asn Phe Pro Val Val Glu Ala Ser Trp Ser

245 250 255 260

15 Met Leu His Asp Glu Arg Pro Tyr Leu Cys Asn Phe Leu Gly Thr Ile

265 270 275

Tyr Glu Asn Ser Ser Arg Gln Ala Leu Met Asn Ile Leu Lys Lys Asp

280 285 290

Gly Asn Asp Lys Leu Cys Trp Val Ser Ala Arg Glu His Trp Gln Pro

20 295 300 305

Gln Glu Thr Asn Glu Ser Leu Lys Asn Tyr Gln Asp Ala Leu Leu Gln

310 315 320

Ser Asp Leu Thr Leu Cys Pro Val Gly Val Asn Thr Glu Cys Tyr Arg

325 330 335 340

25 Ile Tyr Glu Ala Cys Ser Tyr Gly Ser Ile Pro Val Val Glu Asp Val

345 350 355

Met Thr Ala Gly Asn Cys Gly Asn Thr Ser Val His His Gly Ala Pro

360

365

370

Leu Gln Leu Leu Lys Ser Met Gly Ala Pro Phe Ile Phe Ile Lys Asn

375

380

385

Trp Lys Glu Leu Pro Ala Val Leu Glu Lys Glu Lys Thr Ile Ile Leu

5 390

395

400

Gln Glu Lys Ile Glu Arg Arg Lys Met Leu Leu Gln Trp Tyr Gln His

405

410

415

420

Phe Lys Thr Glu Leu Lys Met Lys Phe Thr Asn Ile Leu Glu Ser Ser

425

430

435

10 Phe Leu Met Asn Asn Lys Ser

440

<210> 39

15 <211> 886

<212> DNA

<213> Homo sapiens

<400> 39

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tc atg gag acc ctg ggg gcc ctt ctg gtg ctg gag ttt ctg ctc ctc 107

Met Glu Thr Leu Gly Ala Leu Leu Val Leu Glu Phe Leu Leu Leu

1

5

10

15

tcc ccg gtg gag gcc cag cag gcc acg gag cat cgc ctg aag ccg tgg 155

25 Ser Pro Val Glu Ala Gln Gln Ala Thr Glu His Arg Leu Lys Pro Trp

20

25

30

ctg gtg ggc ctg gct gcg gta gtc ggc ttc ctg atc gtc tat ttg 203

Leu Val Gly Leu Ala Ala Val Val Gly Phe Leu Phe Ile Val Tyr Leu

35 40 45

gtc ttg ctg gcc aac cgc ctc tgg tgt tcc aag gcc agg gct gag gac 251

Val Leu Leu Ala Asn Arg Leu Trp Cys Ser Lys Ala Arg Ala Glu Asp

5 50 55 60

gag gag gag acc acg ttc aga atg gag tcc aac cta tac cag gac cag 299

Glu Glu Glu Thr Thr Phe Arg Met Glu Ser Asn Leu Tyr Gln Asp Gln

65 70 75

agt gaa gac aag aga gag aag aaa gag gcc aag gag aaa gaa gag aag 347

10 Ser Glu Asp Lys Arg Glu Lys Lys Glu Ala Lys Glu Lys Glu Lys

80 85 90 95

agg aag aag gag aaa aag aca gca aag gaa gga gag agc aac ttg gga 395

Arg Lys Lys Glu Lys Lys Thr Ala Lys Glu Gly Glu Ser Asn Leu Gly

100 105 110

15 ctg gat ctg gag gaa aaa gag ccc gga gac cat gag aga gca aag agc 443

Leu Asp Leu Glu Glu Lys Glu Pro Gly Asp His Glu Arg Ala Lys Ser

115 120 125

aca gtc atg tgaagatt cctggctgcc tcttccaggc agtccccag agatgcctct 500

Thr Val Met

20 130

tctgccccctt aaaaggcgtg ccctggactt gaagccgtg aaatgactcc atctggatt 560

cagaatacag tggcttcagaag tgaagaaggc ttggaaaccca ccccacctcc ctatgggg 620

gctctctggg caaacatggt ttcatgcac ccctttctt gagcttggtc cctggctgg 680

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25 ctcatggctc cacctgcttc tccccactgc ctgatttctt ttctctctgc ctgatgtcta 800

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Ser Pro Val Glu Ala Gln Gln Ala Thr Glu His Arg Leu Lys Pro Trp

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35 40 45

15 Val Leu Leu Ala Asn Arg Leu Trp Cys Ser Lys Ala Arg Ala Glu Asp

50 55 60

Glu Glu Glu Thr Thr Phe Arg Met Glu Ser Asn Leu Tyr Gln Asp Gln

65 70 75

Ser Glu Asp Lys Arg Glu Lys Lys Glu Ala Lys Glu Lys Glu Lys

20 80 85 90 95

Arg Lys Lys Glu Lys Lys Thr Ala Lys Glu Gly Glu Ser Asn Leu Gly

100 105 110

Leu Asp Leu Glu Glu Lys Glu Pro Gly Asp His Glu Arg Ala Lys Ser

115 120 125

25 Thr Val Met

130

**DECLARATION, PETITION AND POWER OF ATTORNEY
FOR PATENT APPLICATION**

(Check one):

Declaration Submitted with Initial Filing
 Declaration Submitted after Initial Filing

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

**HUMAN PROTEINS HAVING TRANSMEMBRANE DOMAINS
AND cDNAs ENCODING THESE PROTEINS**

the specification of which (check one):

is attached hereto.
OR
 was filed on **05 October 1998** as PCT International Application Number
**PCT/JP98/04475, and was filed pursuant to 35 U.S.C. §371 as U.S. Serial No.
09/529,205 on April 7, 2000**

and was amended by PCT Article 19 Amendment on _____
(if applicable),

and was amended by PCT Article 34 Amendment on _____
(if applicable).

I acknowledge the duty to disclose to the Office all information known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, §1.56.

I hereby state that I have reviewed and understood the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

PRIORITY CLAIM

(Check one):

no such applications have been filed.
 such applications have been filed as follows

1) FOREIGN PRIORITY CLAIM: I hereby claim foreign priority benefits under Title 35, United States Code, §119(a)-(d) or §365(b) of any foreign application(s) for patent or inventor's certificate or §365(a) of any PCT international application which designated at least one country other than the United States of America, listed below and have also identified below, by checking the box, any foreign application for patent or inventor's certificate or any PCT international application having a filing date before that of the application on which priority is claimed.

Prior Foreign Application Number(s)	Country	Foreign Filing Date (dd/mm/yyyy)	Priority Not Claimed	Certified Copy Attached Yes	Certified Copy Attached No
9/276271	JP	08 October 1997 (08.10.97)	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>
			<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

Additional foreign application numbers are listed on a supplemental priority sheet attached hereto.

2) PROVISIONAL PRIORITY CLAIM: I hereby claim the benefit under Title 35, United States Code §119(e) of any United States provisional application(s) listed below.

Provisional Application Number(s)	Filing Date (dd/mm/yyyy)

Additional provisional application numbers are listed on a supplemental priority sheet attached hereto.

3) U.S./PCT PRIORITY CLAIM: I hereby claim the benefit under Title 35, United States Code, §120 of any United States application or §365(c) of any PCT international application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT international application in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose information which is known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, §1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application.

U.S. Parent Application Number	PCT Parent Number	Parent Filing Date (dd/mm/yyyy)	Parent Patent Number (if applicable)
	PCT/JP98/04475	05 October 1998 (05.10.98)	

Additional U.S. or PCT international application numbers are listed on a supplemental priority sheet attached hereto.

POWER OF ATTORNEY:

As a named inventor, I hereby appoint the following attorneys and/or agents to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.

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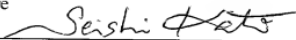
Direct Telephone Calls to: (name and telephone number)

Peter C. Lauro, (617) 227-7400, also at Lahive & Cockfield, LLP, 28 State Street, Boston, Massachusetts 02109, United States of America

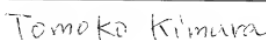
Wherefore I petition that letters patent be granted to me for the invention or discovery described and claimed in the attached specification and claims, and hereby subscribe my name to said specification and claims and to the foregoing declaration, power of attorney, and this petition.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

BD

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